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(54) Title: COMPOSITIONS AND METHODS RELATING TO COLON SPECIFIC GENES AND PROTEINS

(57) Abstract: The present invention relates to newly identified nucleic acid molecules and polypeptides present in normal and neoplastic colon cells, including fragments, variants and derivatives of the nucleic acids and polypeptides. The present invention also relates to antibodies to the polypeptides of the invention, as well as agonists and antagonists of the polypeptides of the invention. The invention also relates to compositions containing the nucleic acid molecules, polypeptides, antibodies, agonists and antagonists of the invention and methods for the use of these compositions. These uses include identifying, diagnosing, monitoring, staging, imaging and treating colon cancer and non-cancerous disease states in colon, identifying colon tissue, monitoring and identifying and/or designing agonists and antagonists of polypeptides of the invention. The uses also include gene therapy, production of transgenic animals and cells, and production of engineered colon tissue for treatment and research.

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COMPOSITIONS AND METHODS RELATING TO COLON SPECIFIC GENES AND PROTEINS

INTRODUCTION

This application claims the benefit of priority from U.S. Provisional Application No. 60/316,259, filed August 31, 2001, which is herein incorporated by reference in its entirety.

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FIELD OF THE INVENTION

The present invention relates to newly identified nucleic acids and polypeptides present in normal and neoplastic colorectal cells, including fragments, variants and derivatives of the nucleic acids and polypeptides. The present invention also relates to antibodies to the polypeptides of the invention, as well as agonists and antagonists of the polypeptides of the invention. The invention also relates to compositions comprising the nucleic acids, polypeptides, antibodies, variants, derivatives, agonists and antagonists of the invention and methods for the use of these compositions. This invention relates to newly developed assays for detecting, diagnosing, monitoring, staging, prognosticating, imaging and treating cancers, particularly colorectal cancer, and identifying and/or designing agonists and antagonists of polypeptides of the invention. The uses also include gene therapy, production of transgenic animals and cells, and production of engineered colorectal tissue for treatment and research.

BACKGROUND OF THE INVENTION

Colorectal cancer is the second most common cause of cancer death in the United States and the third most prevalent cancer in both men and women. M. L. Davila & A. D. 25 Davila, Screening for Colon and Rectal Cancer, in Colon and Rectal Cancer 47 (Peter S. Edelstein ed., 2000). Approximately 100,000 patients every year suffer from colon cancer and approximately half that number die of the disease. Hannah-Ngoc Ha & Bard C. Cosman, Treatment of Colon Cancer, in Colon and Rectal Cancer 157 (Peter S. Edelstein ed., 2000). Nearly all cases of colorectal cancer arise from adenomatous polyps, some of which mature into large polyps, undergo abnormal growth and development, and ultimately progress into cancer. Davila & Davila, supra at 55-56. This progression would appear to take at least 10 years in most patients, rendering it a readily treatable form of

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cancer if diagnosed early, when the cancer is localized. *Id.* at 56; Walter J. Burdette, Cancer: Etiology, Diagnosis, and Treatment 125 (1998).

Although our understanding of the etiology of colon cancer is undergoing continual refinement, extensive research in this area points to a combination of factors, including age, hereditary and nonheriditary conditions, and environmental/dietary factors. Age is a key risk factor in the development of colorectal cancer, Davila & Davila, supra at 48, with men and women over 40 years of age become increasingly susceptible to that cancer, Burdette, supra at 126. Incidence rates increase considerably in each subsequent decade of life. Davila et al., supra at 48. A number of hereditary and nonhereditary 10 conditions have also been linked to a heightened risk of developing colorectal cancer. including familial adenomatous polyposis (FAP), hereditary nonpolyposis colorectal cancer (Lynch syndrome or HNPCC), a personal and/or family history of colorectal cancer or adenomatous polyps, inflammatory bowel disease, diabetes mellitus, and obesity. Id. at 47; Henry T. Lynch & Jane F. Lynch, Hereditary Nonpolyposis Colorectal Cancer (Lynch 15 Syndromes), in Colon and Rectal Cancer 67-68 (Peter S. Edelstein ed., 2000). In the case of FAP, the tumor suppressor gene APC (adenomatous polyposis coli), located at 5q21, has been either mutationally inactivated or deleted. Alberts et al., Molecular Biology of the Cell 1288 (3d ed. 1994). The APC protein plays a role in a number of functions, including cell adhesion, apoptosis, and repression of the c-myc oncogene. N. R. Hall & R. D. Madoff, Genetics and the Polyp-Cancer Sequence, Colon and Rectal Cancer 8 (Peter S. Edelstein, ed., 2000). Of those patients with colorectal cancer who have normal APC genes, over 65% have such mutations in the cancer cells but not in other tissues. Alberts et al., supra at 1288. In the case of HPNCC, patients manifest abnormalities in the tumor suppressor gene HNPCC, but only about 15% of tumors contain the mutated gene. Id. A host of other genes have also been implicated in colorectal cancer, including the K-ras, N-ras, H-ras and c-myc oncogenes, and the tumor suppressor genes DCC (deleted in colon carcinoma) and p53. Hall & Madoff, supra at 8-9; Alberts et al., supra at 1288.

Environmental/dietary factors associated with an increased risk of colorectal cancer include a high fat diet, intake of high dietary red meat, and sedentary lifestyle. Davila & Davila, supra at 47; Reddy, B. S., Prev. Med. 16(4): 460-7 (1987). Conversely, environmental/dietary factors associated with a reduced risk of colorectal cancer include a diet high in fiber, folic acid, calcium, and hormone-replacement therapy in post-

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menopausal women. Davila & Davila, *supra* at 50-55. The effect of antioxidants in reducing the risk of colon cancer is unclear. *Id.* at 53.

Because colon cancer is highly treatable when detected at an early, localized stage, screening should be a part of routine care for all adults starting at age 50, especially those 5 with first-degree relatives with colorectal cancer. One major advantage of colorectal cancer screening over its counterparts in other types of cancer is its ability to not only detect precancerous lesions, but to remove them as well. Davila & Davila, supra at 56. The key colorectal cancer screening tests in use today are fecal occult blood test, sigmoidoscopy, colonoscopy, double-contrast barium enema, and the carcinoembryonic antigen (CEA) test. Id: Burdette, supra at 125.

The fecal occult blood test (FOBT) screens for colorectal cancer by detecting the amount of blood in the stool, the premise being that neoplastic tissue, particularly malignant tissue, bleeds more than typical mucosa, with the amount of bleeding increasing with polyp size and cancer stage. Davila & Davila, supra at 56-57. While effective at 15 detecting early stage tumors, FOBT is unable to detect adenomatous polyps (premalignant lesions), and, depending on the contents of the fecal sample, is subject to rendering false positives. Id. at 56-59. Sigmoidoscopy and colonoscopy, by contrast, allow direct visualization of the bowel, and enable one to detect, biopsy, and remove adenomatous polyps. Id. at 59-60, 61. Despite the advantages of these procedures, there are accompanying downsides: sigmoidoscopy, by definition, is limited to the sigmoid colon and below, colonoscopy is a relatively expensive procedure, and both share the risk of possible bowel perforation and hemorrhaging. Id. at 59-60. Double-contrast barium enema (DCBE) enables detection of lesions better than FOBT, and almost as well a colonoscopy, but it may be limited in evaluating the winding rectosigmoid region. Id. at 25 60. The CEA blood test, which involves screening the blood for carcinoembryonic antigen, shares the downside of FOBT, in that it is of limited utility in detecting colorectal cancer at an early stage. Burdette, supra at 125.

Once colon cancer has been diagnosed, treatment decisions are typically made in reference to the stage of cancer progression. A number of techniques are employed to stage the cancer (some of which are also used to screen for colon cancer), including pathologic examination of resected colon, sigmoidoscopy, colonoscopy, and various imaging techniques. <u>AJCC Cancer Staging Handbook</u> 84 (Irvin D. Fleming et al. eds., 5th ed. 1998). Montgomery, R. C. and Ridge, J.A., Semin. Surg. Oncol. 15(3): 143-150

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(1998). Moreover, chest films, liver functionality tests, and liver scans are employed to determine the extent of metastasis. Fleming et al. eds., supra at 84. While computerized tomography and magnetic resonance imaging are useful in staging colorectal cancer in its later stages, both have unacceptably low staging accuracy for identifying early stages of the disease, due to the difficulty that both methods have in (1) revealing the depth of bowel wall tumor infiltration and (2) diagnosing malignant adenopathy. Thoeni, R. F., Radiol. Clin. N. Am. 35(2): 457-85 (1997). Rather, techniques such as transrectal ultrasound (TRUS) are preferred in this context, although this technique is inaccurate with respect to detecting small lymph nodes that may contain metastases. David Blumberg & Frank G. Opelka, Neoadjuvant and Adjuvant Therapy for Adenocarcinoma of the Rectum, in Colon and Rectal Cancer 316 (Peter S. Edelstein ed., 2000).

Several classification systems have been devised to stage the extent of colorectal cancer, including the Dukes' system and the more detailed International Union against Cancer-American Joint Committee on Cancer TNM staging system, which is considered by many in the field to be a more useful staging system. Burdette, supra at 126-27. The TNM system, which is used for either clinical or pathological staging, is divided into four stages, each of which evaluates the extent of cancer growth with respect to primary tumor (T), regional lymph nodes (N), and distant metastasis (M). Fleming et al. eds., supra at 84-85. The system focuses on the extent of tumor invasion into the intestinal wall, invasion of adjacent structures, the number of regional lymph nodes that have been affected, and whether distant metastasis has occurred. Id. at 81.

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Stage 0 is characterized by in situ carcinoma (Tis), in which the cancer cells are located inside the glandular basement membrane (intraepithelial) or lamina propria (intramucosal). Id. at 84-85; Burdette, supra at 127. In this stage, the cancer has not spread to the regional lymph nodes (N0), and there is no distant metastasis (M0). Fleming et al. eds., supra at 85; Burdette, supra at 127. In stage I, there is still no spread of the cancer to the regional lymph nodes and no distant metastasis, but the tumor has invaded the submucosa (T1) or has progressed further to invade the muscularis propria (T2). Fleming et al. eds., supra at 84-85; Burdette, supra at 127. Stage II also involves no spread of the cancer to the regional lymph nodes and no distant metastasis, but the tumor has invaded the subserosa, or the nonperitonealized pericolic or perirectal tissues (T3), or has progressed to invade other organs or structures, and/or has perforated the visceral peritoneum (T4). Id. Stage 3 is characterized by any of the T substages, no distant

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metastasis, and either metastasis in 1 to 3 regional lymph nodes (N1) or metastasis in four or more regional lymph nodes (N2). Fleming et al. eds., supra at 85; Burdette, supra at 127. Lastly, stage 4 involves any of the T or N substages, as well as distant metastasis. Id.

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Currently, pathological staging of colon cancer is preferable over clinical staging as pathological staging provides a more accurate prognosis. Pathological staging typically involves examination of the resected colon section, along with surgical examination of the abdominal cavity. Fleming et al. eds., supra at 84. Clinical staging would be a preferred method of staging were it at least as accurate as pathological staging, as it does not depend on the invasive procedures of its counterpart.

Turning to the treatment of colorectal cancer, surgical resection results in a cure for roughly 50% of patients. Burdette, supra at 12S. Irradiation is used both preoperatively and postoperatively in treating colorectal cancer. Id. at 12S, 132-33. Chemotherapeutic agents, particularly 5-fluorouracil, are also powerful weapons in treating colorectal cancer. Id. at 12S, 133. Other agents include irinotecan and floxuridine, cisplatin, levamisole, methotrexate, interferon-α, and leucovorin. Id. at 133. Nonetheless, thirty to forty percent of patients will develop a recurrence of colon cancer following surgical resection. Wayne De Vos, Follow-up After Treatment of Colon Cancer, Colon and Rectal Cancer 225 (Peter S. Edelstein ed., 2000), which in many patients is the ultimate cause of death.

Accordingly, colon cancer patients must be closely monitored to determine response to therapy and to detect persistent or recurrent disease and metastasis.
From the foregoing, it is clear that procedures used for detecting, diagnosing, monitoring, staging, prognosticating, and preventing the recurrence of colorectal cancer are of critical importance to the outcome of the patient. Moreover, current procedures, while helpful in each of these analyses, are limited by their specificity, sensitivity, invasiveness, and/or their cost. As such, highly specific and sensitive procedures that would operate by way of detecting novel markers in cells, tissues, or bodily fluids, with minimal invasiveness and at a reasonable cost, would be highly desirable.

Accordingly, there is a great need for more sensitive and accurate methods for predicting whether a person is likely to develop colorectal cancer, for diagnosing colorectal cancer, for monitoring the progression of the disease, for staging the colorectal cancer, for determining whether the colorectal cancer has metastasized, and for imaging the colorectal cancer. There is also a need for better treatment of colorectal cancer.

6 SUMMARY OF THE INVENTION

The present invention solves many needs in the art by providing nucleic acid molecules, polypeptides and antibodies thereto, variants and derivatives of the nucleic acids and polypeptides, agonists and antagonists that may be used to identify, diagnose, monitor, stage, image and treat colon cancer and non-cancerous disease states in colon; identify and monitor colon tissue; and identify and design agonists and antagonists of polypeptides of the invention. The invention also provides gene therapy, methods for producing transgenic animals and cells, and methods for producing engineered colon tissue for treatment and research.

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One aspect of the present invention relates to nucleic acid molecules that are specific to colon cells, colon tissue and/or the colon organ. These colon specific nucleic acids (CSNAs) may be a naturally-occurring cDNA, genomic DNA, RNA, or a fragment of one of these nucleic acids, or may be a non-naturally-occurring nucleic acid molecule. If the CSNA is genomic DNA, then the CSNA is a colon specific gene (CSG). In a preferred embodiment, the nucleic acid molecule encodes a polypeptide that is specific to colon. More preferred is a nucleic acid molecule encodes a polypeptide comprising an amino acid sequence of SEQ ID NO: 101-190. In another preferred embodiment, the nucleic acid molecule comprises a nucleic acid sequence of SEQ ID NO: 1-100. For the sequences listed herein, DEX0234_101 corresponds to SEQ ID NO: 1, DEX0234_101 corresponds to SEQ ID NO: 101, etc.

This aspect of the present invention also relates to nucleic acid molecules that selectively hybridize or exhibit substantial sequence similarity to nucleic acid molecules encoding a Colon Specific Protein (CSP), or that selectively hybridize or exhibit substantial sequence similarity to a CSNA. In one embodiment of the present invention the nucleic acid molecule comprises an allelic variant of a nucleic acid molecule encoding a CSP, or an allelic variant of a CSNA. In another embodiment, the nucleic acid molecule comprises a part of a nucleic acid sequence that encodes a CSP or a part of a nucleic acid sequence of a CSNA.

In addition, this aspect of the present invention relates to a nucleic acid molecule further comprising one or more expression control sequences controlling the transcription and/or translation of all or a part of a CSNA or the the transcription and/or translation of a nucleic acid molecule that encodes all or a fragment of a CSP.

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Another aspect of the present invention relates to vectors and/or host cells comprising a nucleic acid molecule of this invention. In a preferred embodiment, the nucleic acid molecule of the vector and/or host cell encodes all or a fragment of a CSP. In another preferred embodiment, the nucleic acid molecule of the vector and/or host cell comprises all or a part of a CSNA. Vectors and host cells of the present invention are useful in the recombinant production of polypeptides, particularly CSPs of the present invention.

Another aspect of the present invention relates to polypeptides encoded by a nucleic acid molecule of this invention. The polypeptide may comprise either a fragment or a full-length protein. In a preferred embodiment, the polypeptide is a CSP. However, this aspect of the present invention also relates to mutant proteins (muteins) of CSPs, fusion proteins of which a portion is a CSP, and proteins and polypeptides encoded by allelic variants of a CSNA as provided herein.

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Another aspect of the present invention relates to antibodies and other binders that specifically binds to a polypeptide of the instant invention. Accordingly antibodies or binders of the present specifically bind to CSPs, muteins, fusion proteins, and/or homologous proteins or a polypeptides encoded by allelie variants of an CSNA as provided herein.

Another aspect of the present invention relates to agonists and antagonists of the nucleic acid molecules and polypeptides of this invention. The agonists and antagonists of the instant invention may be used to treat colon cancer and non-cancerous disease states in colon and to produce engineered colon tissue.

Another aspect of the present invention relates to methods for using the nucleic acid molecules to detect or amplify nucleic acid molecules that have similar or identical nucleic acid sequences compared to the nucleic acid molecules described herein. Such methods are useful in identifying, diagnosing, monitoring, staging, imaging and treating colon cancer and non-cancerous disease states in colon. Such methods are also useful in identifying and/or monitoring colon tissue. In addition, measurement of levels of the nucleic acid molecules of this invention may be useful for diagnostics as part of panel in combination with other markers.

Another aspect of the present invention relates to use of the nucleic acid molecules of this invention in gene therapy, for producing transgenic animals and cells, and for producing engineered colon tissue for treatment and research.

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Another aspect of the present invention relates to methods for detecting polypeptides this invention, preferably using antibodies thereto. Such methods are useful to identify, diagnose, monitor, stage, image and treat colon cancer and non-cancerous disease states in colon. In addition, measurement of levels of the polypeptides of this invention may be useful to identify, diagnose, monitor, stage, image colon cancer in combination with other colon cancer markers. The polypeptides of the present invention can also be used to identify and/or monitor colon tissue, and to produce engineered colon tissue.

Yet another aspect of the present invention relates to a computer readable means of storing the nucleic acid and amino acid sequences of the invention. The records of the computer readable means can be accessed for reading and displaying of sequences for comparison, alignment and ordering of the sequences of the invention to other sequences. In addition, the computer records regarding the nucleic acid and/or amino acid sequences and/or measurements of their levels may be used alone or in combination with other markers to diagnose colon related diseases.

DETAILED DESCRIPTION OF THE INVENTION

Definitions and General Techniques

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Unless otherwise defined herein, scientific and technical terms used in connection 20 with the present invention shall have the meanings that are commonly understood by those of ordinary skill in the art. Further, unless otherwise required by context, singular terms shall include pluralities and plural terms shall include the singular. Generally, nomenclatures used in connection with, and techniques of, cell and tissue culture, molecular biology, immunology, microbiology, genetics and protein and nucleic acid chemistry and hybridization described herein are those well-known and commonly used in 25 the art. The methods and techniques of the present invention are generally performed according to conventional methods well-known in the art and as described in various general and more specific references that are cited and discussed throughout the present specification unless otherwise indicated. See, e.g., Sambrook et al., Molecular Cloning: A Laboratory Manual, 2d ed., Cold Spring Harbor Laboratory Press (1989) and Sambrook et al., Molecular Cloning: A Laboratory Manual, 3d ed., Cold Spring Harbor Press (2001): Ausubel et al., Current Protocols in Molecular Biology, Greene Publishing Associates (1992, and Supplements to 2000): Ausubel et al., Short Protocols in Molecular Biology: A

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Compendium of Methods from Current Protocols in Molecular Biology – 4th Ed., Wiley & Sons (1999); Harlow and Lane, Antibodies: A Laboratory Manual, Cold Spring Harbor Laboratory Press (1990); and Harlow and Lane, Using Antibodies: A Laboratory Manual, Cold Spring Harbor Laboratory Press (1999).

Enzymatic reactions and purification techniques are performed according to manufacturer's specifications, as commonly accomplished in the art or as described herein. The nomenclatures used in connection with, and the laboratory procedures and techniques of, analytical chemistry, synthetic organic chemistry, and medicinal and pharmaceutical chemistry described herein are those well-known and commonly used in the art. Standard techniques are used for chemical syntheses, chemical analyses, pharmaceutical preparation, formulation, and delivery, and treatment of patients.

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The following terms, unless otherwise indicated, shall be understood to have the following meanings:

A "nucleic acid molecule" of this invention refers to a polymeric form of nucleotides and includes both sense and antisense strands of RNA, cDNA, genomic DNA, and synthetic forms and mixed polymers of the above. A nucleotide refers to a ribonucleotide, deoxynucleotide or a modified form of either type of nucleotide. A "nucleic acid molecule" as used herein is synonymous with "nucleic acid" and "polynucleotide." The term "nucleic acid molecule" usually refers to a molecule of at least 10 bases in length, unless otherwise specified. The term includes single and double stranded forms of DNA. In addition, a polynucleotide may include either or both naturally-occurring and modified nucleotides linked together by naturally-occurring and/or non-naturally occurring nucleotide linkages.

The nucleic acid molecules may be modified chemically or biochemically or may contain non-natural or derivatized nucleotide bases, as will be readily appreciated by those of skill in the art. Such modifications include, for example, labels, methylation, substitution of one or more of the naturally occurring nucleotides with an analog, internucleotide modifications such as uncharged linkages (e.g., methyl phosphonates, phosphoramidates, carbamates, etc.), charged linkages (e.g., phosphorothioates, phosphorodithioates, etc.), pendent moieties (e.g., polypeptides), intercalators (e.g., acridine, psoralen, etc.), chelators, alkylators, and modified linkages (e.g., alpha anomeric nucleic acids, etc.) The term "nucleic acid molecule" also includes any topological conformation, including single-stranded, double-stranded, partially

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duplexed, triplexed, hairpinned, circular and padlocked conformations. Also included are synthetic molecules that mimic polynucleotides in their ability to bind to a designated sequence via hydrogen bonding and other chemical interactions. Such molecules are known in the art and include, for example, those in which peptide linkages substitute for phosphate linkages in the backbone of the molecule.

A "gene" is defined as a nucleic acid molecule that comprises a nucleic acid sequence that encodes a polypeptide and the expression control sequences that surround the nucleic acid sequence that encodes the polypeptide. For instance, a gene may comprise a promoter, one or more enhancers, a nucleic acid sequence that encodes a polypeptide, downstream regulatory sequences and, possibly, other nucleic acid sequences involved in regulation of the expression of an RNA. As is well-known in the art. eukarvotic genes usually contain both exons and introns. The term "exon" refers to a nucleic acid sequence found in genomic DNA that is bioinformatically predicted and/or experimentally confirmed to contribute contiguous sequence to a mature mRNA 15 transcript. The term "intron" refers to a nucleic acid sequence found in genomic DNA that is predicted and/or confirmed to not contribute to a mature mRNA transcript, but rather to be "spliced out" during processing of the transcript.

A nucleic acid molecule or polypeptide is "derived" from a particular species if the nucleic acid molecule or polypeptide has been isolated from the particular species, or if the nucleic acid molecule or polypentide is homologous to a nucleic acid molecule or polypeptide isolated from a particular species.

An "isolated" or "substantially pure" nucleic acid or polynucleotide (e.g., an RNA, DNA or a mixed polymer) is one which is substantially separated from other cellular components that naturally accompany the native polynucleotide in its natural host cell. 25 e.g., ribosomes, polymerases, or genomic sequences with which it is naturally associated. The term embraces a nucleic acid or polynucleotide that (1) has been removed from its naturally occurring environment, (2) is not associated with all or a portion of a polynucleotide in which the "isolated polynucleotide" is found in nature, (3) is operatively linked to a polynucleotide which it is not linked to in nature. (4) does not occur in nature as part of a larger sequence or (5) includes nucleotides or internucleoside bonds that are not found in nature. The term "isolated" or "substantially pure" also can be used in reference to recombinant or cloned DNA isolates, chemically synthesized polynucleotide analogs, or polynucleotide analogs that are biologically synthesized by heterologous

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systems. The term "isolated nucleic acid molecule" includes nucleic acid molecules that are integrated into a host cell chromosome at a heterologous site, recombinant fusions of a native fragment to a heterologous sequence, recombinant vectors present as episomes or as integrated into a host cell chromosome.

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A "part" of a nucleic acid molecule refers to a nucleic acid molecule that comprises a partial contiguous sequence of at least 10 bases of the reference nucleic acid molecule. Preferably, a part comprises at least 15 to 20 bases of a reference nucleic acid molecule. In theory, a nucleic acid sequence of 17 nucleotides is of sufficient length to occur at random less frequently than once in the three gigabase human genome, and thus to provide a nucleic acid probe that can uniquely identify the reference sequence in a nucleic acid mixture of genomic complexity. A preferred part is one that comprises a nucleic acid sequence that can encode at least 6 contiguous amino acid sequences (fragments of at least 18 nucleotides) because they are useful in directing the expression or synthesis of peptides that are useful in mapping the epitopes of the polypeptide encoded 15 by the reference nucleic acid. See, e.g., Gevsen et al., Proc. Natl. Acad. Sci. USA 81:3998-4002 (1984); and United States Patent Nos. 4,708,871 and 5,595,915, the disclosures of which are incorporated herein by reference in their entireties. A part may also comprise at least 25, 30, 35 or 40 nucleotides of a reference nucleic acid molecule, or at least 50, 60, 70, 80, 90, 100, 150, 200, 250, 300, 350, 400 or 500 nucleotides of a reference nucleic acid molecule. A part of a nucleic acid molecule may comprise no other nucleic acid sequences. Alternatively, a part of a nucleic acid may comprise other nucleic acid sequences from other nucleic acid molecules.

The term "oligonucleotide" refers to a nucleic acid molecule generally comprising a length of 200 bases or fewer. The term often refers to single-stranded 25 deoxyribonucleotides, but it can refer as well to single-or double-stranded ribonucleotides. RNA: DNA hybrids and double-stranded DNAs, among others, Preferably, oligonucleotides are 10 to 60 bases in length and most preferably 12, 13, 14, 15, 16, 17, 18, 19 or 20 bases in length. Other preferred oligonucleotides are 25, 30, 35, 40, 45, 50, 55 or 60 bases in length. Oligonucleotides may be single-stranded, e.g. for use as probes or primers, or may be double-stranded, e.g. for use in the construction of a mutant gene. Oligonucleotides of the invention can be either sense or antisense oligonucleotides. An oligonucleotide can be derivatized or modified as discussed above for nucleic acid molecules.

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Oligonucleotides, such as single-stranded DNA probe oligonucleotides, often are synthesized by chemical methods, such as those implemented on automated oligonucleotide synthesizers. However, oligonucleotides can be made by a variety of other methods, including in vitro recombinant DNA-mediated techniques and by

5 expression of DNAs in cells and organisms. Initially, chemically synthesized DNAs typically are obtained without a 5' phosphate. The 5' ends of such oligonucleotides are not substrates for phosphodiester bond formation by ligation reactions that employ DNA ligases typically used to form recombinant DNA molecules. Where ligation of such oligonucleotides is desired, a phosphate can be added by standard techniques, such as those that employ a kinase and ATP. The 3' end of a chemically synthesized oligonucleotide generally has a free hydroxyl group and, in the presence of a ligase, such as T4 DNA ligase, readily will form a phosphodiester bond with a 5' phosphate of another polynucleotide, such as another oligonucleotide. As is well-known, this reaction can be prevented selectively, where desired, by removing the 5' phosphates of the other

The term "naturally-occurring nucleotide" referred to herein includes naturally-occurring deoxyribonucleotides and ribonucleotides. The term "modified nucleotides" referred to herein includes nucleotides with modified or substituted sugar groups and the like. The term "nucleotide linkages" referred to herein includes nucleotides linkages such as phosphorothioate, phosphorodithioate, phosphoroanilothioate, phosphoroaniladate, phosphoroanidate, and the like. See e.g., LaPlanche et al. Nucl. Acids Res. 14:9081-9093 (1986); Stein et al. Nucl. Acids Res. 16:3209-3221 (1988); Zon et al. Anti-Cancer Drug Design 6:539-568 (1991); Zon et al., in Eckstein (ed.) Oligonucleotides and Analogues: A Practical Approach, pp. 87-108, 25 Oxford University Press (1991); Uhlmann and Peyman Chemical Reviews 90:543 (1990), and United States Patent No. 5,151,510, the disclosure of which is hereby incorporated by reference in its entirety.

Unless specified otherwise, the left hand end of a polynucleotide sequence in sense orientation is the 5' end and the right hand end of the sequence is the 3' end. In addition, the left hand direction of a polynucleotide sequence in sense orientation is referred to as the 5' direction, while the right hand direction of the polynucleotide sequence is referred to as the 3' direction. Further, unless otherwise indicated, each nucleotide sequence is set forth herein as a sequence of deoxyribonucleotides. It is intended, however, that the given

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sequence be interpreted as would be appropriate to the polynucleotide composition: for example, if the isolated nucleic acid is composed of RNA, the given sequence intends ribonucleotides, with uridine substituted for thymidine.

The term "allelic variant" refers to one of two or more alternative naturallyoccurring forms of a gene, wherein each gene possesses a unique nucleotide sequence. In a preferred embodiment, different alleles of a given gene have similar or identical biological properties.

The term "percent sequence identity" in the context of nucleic acid sequences refers to the residues in two sequences which are the same when aligned for maximum 10 correspondence. The length of sequence identity comparison may be over a stretch of at least about nine nucleotides, usually at least about 20 nucleotides, more usually at least about 24 nucleotides, typically at least about 28 nucleotides, more typically at least about 32 nucleotides, and preferably at least about 36 or more nucleotides. There are a number of different algorithms known in the art which can be used to measure nucleotide sequence 15 identity. For instance, polynucleotide sequences can be compared using FASTA, Gap or Bestfit, which are programs in Wisconsin Package Version 10.0, Genetics Computer Group (GCG), Madison, Wisconsin. FASTA, which includes, e.g., the programs FASTA2 and FASTA3, provides alignments and percent sequence identity of the regions of the best overlap between the query and search sequences (Pearson, Methods Enzymol. 183: 63-98 (1990): Pearson, Methods Mol. Biol. 132: 185-219 (2000): Pearson, Methods Enzymol. 266: 227-258 (1996); Pearson, J. Mol. Biol. 276: 71-84 (1998)). Unless otherwise specified, default parameters for a particular program or algorithm are used. For instance, percent sequence identity between nucleic acid sequences can be determined using FASTA with its default parameters (a word size of 6 and the NOPAM factor for the scoring matrix) or using Gap with its default parameters as provided in GCG Version 6.1.

A reference to a nucleic acid sequence encompasses its complement unless otherwise specified. Thus, a reference to a nucleic acid molecule having a particular sequence should be understood to encompass its complementary strand, with its complementary sequence. The complementary strand is also useful, e.g., for antisense therapy, hybridization probes and PCR primers.

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In the molecular biology art, researchers use the terms "percent sequence identity", "percent sequence similarity" and "percent sequence homology" interchangeably. In this

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application, these terms shall have the same meaning with respect to nucleic acid sequences only.

The term "substantial similarity" or "substantial sequence similarity," when referring to a nucleic acid or fragment thereof, indicates that, when optimally aligned with appropriate nucleotide insertions or deletions with another nucleic acid (or its complementary strand), there is nucleotide sequence identity in at least about 50%, more preferably 60% of the nucleotide bases, usually at least about 70%, more usually at least about 80%, preferably at least about 90%, and more preferably at least about 95-98% of the nucleotide bases, as measured by any well-known algorithm of sequence identity, such 10 as FASTA, BLAST or Gap, as discussed above.

Alternatively, substantial similarity exists between a first and second nucleic acid sequence when the first nucleic acid sequence or fragment thereof hybridizes to an antisense strand of the second nucleic acid, , under selective hybridization conditions. Typically, selective hybridization will occur between the first nucleic acid sequence and an antisense strand of the second nucleic acid sequence when there is at least about 55% sequence identity between the first and second nucleic acid sequences-preferably at least about 65%, more preferably at least about 75%, and most preferably at least about 90% over a stretch of at least about 14 nucleotides, more preferably at least 17 nucleotides, even more preferably at least 20, 25, 30, 35, 40, 50, 60, 70, 80, 90 or 100 nucleotides.

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Nucleic acid hybridization will be affected by such conditions as salt concentration, temperature, solvents, the base composition of the hybridizing species, length of the complementary regions, and the number of nucleotide base mismatches between the hybridizing nucleic acids, as will be readily appreciated by those skilled in the art. "Stringent hybridization conditions" and "stringent wash conditions" in the context 25 of nucleic acid hybridization experiments depend upon a number of different physical parameters. The most important parameters include temperature of hybridization, base composition of the nucleic acids, salt concentration and length of the nucleic acid. One having ordinary skill in the art knows how to vary these parameters to achieve a particular stringency of hybridization. In general, "stringent hybridization" is performed at about 25°C below the thermal melting point (Tm) for the specific DNA hybrid under a particular set of conditions. "Stringent washing" is performed at temperatures about 5°C lower than the Tm for the specific DNA hybrid under a particular set of conditions. The Tm is the

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temperature at which 50% of the target sequence hybridizes to a perfectly matched probe.
See Sambrook (1989), supra, p. 9.51.

The T_m for a particular DNA-DNA hybrid can be estimated by the formula:

 $T_m = 81.5^{\circ}C + 16.6 (\log_{10}[Na^+]) + 0.41 (fraction G + C) -$

5 0.63 (% formamide) - (600/l) where l is the length of the hybrid in base pairs.

The T_m for a particular RNA-RNA hybrid can be estimated by the formula;

 $T_m = 79.8^{\circ}C + 18.5 (\log_{10}[Na^+]) + 0.58 (fraction G + C) +$

11.8 (fraction G + C)2 - 0.35 (% formamide) - (820/l).

The T_m for a particular RNA-DNA hybrid can be estimated by the formula:

10 $T_m = 79.8^{\circ}C + 18.5(log_{10}[Na^+]) + 0.58 \text{ (fraction G + C)} +$

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11.8 (fraction G + C)² - 0.50 (% formamide) - (820/1).

In general, the T_m decreases by 1-1.5°C for each 1% of mismatch between two nucleic acid sequences. Thus, one having ordinary skill in the art can alter hybridization and/or washing conditions to obtain sequences that have higher or lower degrees of sequence identity to the target nucleic acid. For instance, to obtain hybridizing nucleic acids that contain up to 10% mismatch from the target nucleic acid sequence, 10-15°C would be subtracted from the calculated T_m of a perfectly matched hybrid, and then the hybridization and washing temperatures adjusted accordingly. Probe sequences may also hybridize specifically to duplex DNA under certain conditions to form triplex or other

higher order DNA complexes. The preparation of such probes and suitable hybridization conditions are well-known in the art.

An example of stringent hybridization conditions for hybridization of complementary nucleic acid sequences having more than 100 complementary residues on a filter in a Southern or Northern blot or for screening a library is 50% formamide/6X SSC at 42°C for at least ten hours and preferably overnight (approximately 16 hours). Another example of stringent hybridization conditions is 6X SSC at 68°C without formamide for at least ten hours and preferably overnight. An example of moderate stringency hybridization conditions is 6X SSC at 55°C without formamide for at least ten hours and preferably overnight. An example of low stringency hybridization conditions for

60 hybridization of complementary nucleic acid sequences having more than 100 complementary residues on a filter in a Southern or northern blot or for screening a library is 6X SSC at 42°C for at least ten hours. Hybridization conditions to identify nucleic acid sequences that are similar but not identical can be identified by experimentally changing

the hybridization temperature from 68°C to 42°C while keeping the salt concentration constant (6X SSC), or keeping the hybridization temperature and salt concentration constant (e.g. 42°C and 6X SSC) and varying the formamide concentration from 50% to 0%. Hybridization buffers may also include blocking agents to lower background. These agents are well-known in the art. See Sambrook et al. (1989), supra, pages 8.46 and 9.46-9.58. See also Ausubel (1992), supra, Ausubel (1999), supra, and Sambrook (2001), supra

Wash conditions also can be altered to change stringency conditions. An example of stringent wash conditions is a 0.2x SSC wash at 65°C for 15 minutes (see Sambrook (1989), supra, for SSC buffer). Often the high stringency wash is preceded by a low stringency wash to remove excess probe. An exemplary medium stringency wash for duplex DNA of more than 100 base pairs is 1x SSC at 45°C for 15 minutes. An exemplary low stringency wash for such a duplex is 4x SSC at 40°C for 15 minutes. In general, signal-to-noise ratio of 2x or higher than that observed for an unrelated probe in the particular hybridization assay indicates detection of a specific hybridization.

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As defined herein, nucleic acids that do not hybridize to each other under stringent conditions are still substantially similar to one another if they encode polypeptides that are substantially identical to each other. This occurs, for example, when a nucleic acid is created synthetically or recombinantly using a high codon degeneracy as permitted by the redundancy of the genetic code.

Hybridization conditions for nucleic acid molecules that are shorter than 100 nucleotides in length (e.g., for oligonucleotide probes) may be calculated by the formula:

 $T_m = 81.5^{\circ}C + 16.6(log_{10}[Na^{\dagger}]) + 0.41(fraction G+C) - (600/N)$, wherein N is change length and the [Na[†]] is 1 M or less. See Sambrook (1989), supra, p. 11.46. For hybridization of probes shorter than 100 nucleotides, hybridization is usually performed under stringent conditions (5-10°C below the T_m) using high concentrations (0.1-1.0 pmol/ml) of probe. Id. at p. 11.45. Determination of hybridization using mismatched probes, pools of degenerate probes or "guessmers," as well as hybridization solutions and methods for empirically determining hybridization conditions are well-known in the art. See, e.g., Ausubel (1999), supra; Sambrook (1989), supra, pp. 11.45-11.57.

The term "digestion" or "digestion of DNA" refers to catalytic cleavage of the DNA with a restriction enzyme that acts only at certain sequences in the DNA. The various restriction enzymes referred to herein are commercially available and their

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reaction conditions, cofactors and other requirements for use are known and routine to the skilled artisan. For analytical purposes, typically, 1 µg of plasmid or DNA fragment is digested with about 2 units of enzyme in about 20 µl of reaction buffer. For the purpose of isolating DNA fragments for plasmid construction, typically 5 to 50 µg of DNA are

5 digested with 20 to 250 units of enzyme in proportionately larger volumes. Appropriate buffers and substrate amounts for particular restriction enzymes are described in standard laboratory manuals, such as those referenced below, and they are specified by commercial suppliers. Incubation times of about 1 hour at 37°C are ordinarily used, but conditions may vary in accordance with standard procedures, the supplier's instructions and the particulars of the reaction. After digestion, reactions may be analyzed, and fragments may be purified by electrophoresis through an agarose or polyacrylamide gel, using well-known methods that are routine for those skilled in the art.

The term "ligation" refers to the process of forming phosphodiester bonds between two or more polynucleotides, which most often are double-stranded DNAs. Techniques for ligation are well-known to the art and protocols for ligation are described in standard laboratory manuals and references, such as, e.g., Sambrook (1989), supra.

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Genome-derived "single exon probes," are probes that comprise at least part of an exon ("reference exon") and can hybridize detectably under high stringency conditions to transcript-derived nucleic acids that include the reference exon but do not hybridize detectably under high stringency conditions to nucleic acids that lack the reference exon. Single exon probes typically further comprise, contiguous to a first end of the exon portion, a first intronic and/or intergenic sequence that is identically contiguous to the exon in the genome, and may contain a second intronic and/or intergenic sequence that is identically contiguous to the exon in the genome. The minimum length of genomederived single exon probes is defined by the requirement that the exonic portion be of sufficient length to hybridize under high stringency conditions to transcript-derived nucleic acids, as discussed above. The maximum length of genome-derived single exon probes is defined by the requirement that the probes contain portions of no more than one exon. The single exon probes may contain priming sequences not found in contiguity with the rest of the probe sequence in the genome, which priming sequences are useful for PCR and other amplification-based technologies. In another aspect, the invention is directed to single exon probes based on the CSNAs disclosed herein.

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In one embodiment, the term "microarray" refers to a "nucleic acid microarray" having a substrate-bound plurality of nucleic acids, hybridization to each of the plurality of bound nucleic acids being separately detectable. The substrate can be solid or porous. planar or non-planar, unitary or distributed. Nucleic acid microarrays include all the devices so called in Schena (ed.), DNA Microarrays: A Practical Approach (Practical Approach Series), Oxford University Press (1999); Nature Genet. 21(1)(suppl.):1 - 60 (1999); Schena (ed.), Microarray Biochip: Tools and Technology, Eaton Publishing Company/BioTechniques Books Division (2000). Additionally, these nucleic acid microarrays include substrate-bound plurality of nucleic acids in which the plurality of nucleic acids are disposed on a plurality of beads, rather than on a unitary planar substrate, as is described, inter alia, in Brenner et al., Proc. Natl. Acad. Sci. USA 97(4):1665-1670 (2000). Examples of nucleic acid microarrays may be found in U.S. Patent Nos. 6,391,623, 6,383,754, 6,383,749, 6,380,377, 6,379,897, 6,376,191, 6,372,431, 6,351,712 6,344,316, 6,316,193, 6,312,906, 6,309,828, 6,309,824, 6,306,643, 6,300,063, 6,287,850, 6,284,497, 6,284,465, 6,280,954, 6,262,216, 6,251,601, 6,245,518, 6,263,287, 6,251,601, 6,238,866, 6,228,575, 6,214,587, 6,203,989, 6,171,797, 6,103,474, 6,083,726, 6,054,274, 6,040,138, 6,083,726, 6,004,755, 6,001,309, 5,958,342, 5,952,180, 5,936,731, 5,843,655, 5,814,454, 5,837,196, 5,436,327, 5,412,087, 5,405,783, the disclosures of which are

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In an alternative embodiment, a "microarray" may also refer to a "peptide microarray" or "protein microarray" having a substrate-bound collection of plurality of polypeptides, the binding to each of the plurality of bound polypeptides being separately detectable. Alternatively, the peptide microarray, may have a plurality of binders, including but not limited to monoclonal antibodies, polyclonal antibodies, phage display binders, yeast 2 hybrid binders, aptamers, which can specifically detect the binding of the polypeptides of this invention. The array may be based on autoantibody detection to the polypeptides of this invention, see Robinson et al., Nature Medicine 8(3):295-301 (2002).

Examples of peptide arrays may be found in WO 02/31463, WO 02/25288, WO 01/94946.

incorporated herein by reference in their entireties.

WO 01/88162, WO 01/68671, WO 01/57259, WO 00/61806, WO 00/54046, WO 00/47774, WO 99/40434, WO 99/39210, WO 97/42507 and U.S. Patent Nos. 6,268,210, 5,766,960, 5,143,854, the disclosures of which are incorporated herein by reference in their entirelies.

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In addition, determination of the levels of the CSNA or CSP may be made in a multiplex manner using techniques described in WO 02/29109, WO 02/24959, WO 01/83502, WO01/73113, WO 01/59432, WO 01/57269, WO 99/67641, the disclosures of which are incorporated herein by reference in their entireties.

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The term "mutant", "mutated", or "mutation" when applied to nucleic acid sequences means that nucleotides in a nucleic acid sequence may be inserted, deleted or changed compared to a reference nucleic acid sequence. A single alteration may be made at a locus (a point mutation) or multiple nucleotides may be inserted, deleted or changed at a single locus. In addition, one or more alterations may be made at any number of loci within a nucleic acid sequence. In a preferred embodiment of the present invention, the nucleic acid sequence is the wild type nucleic acid sequence encoding a CSP or is a CSNA. The nucleic acid sequence may be mutated by any method known in the art including those mutagenesis techniques described infra.

The term "error-prone PCR" refers to a process for performing PCR under conditions where the copying fidelity of the DNA polymerase is low, such that a high rate of point mutations is obtained along the entire length of the PCR product. See, e.g., Leung et al., Technique 1: 11-15 (1989) and Caldwell et al., PCR Methods Applic. 2: 28-33 (1992).

The term "oligonucleotide-directed mutagenesis" refers to a process which enables the generation of site-specific mutations in any cloned DNA segment of interest. See, e.g., Reidhaar-Olson et al., Science 241: 53-57 (1988).

The term "assembly PCR" refers to a process which involves the assembly of a PCR product from a mixture of small DNA fragments. A large number of different PCR reactions occur in parallel in the same vial, with the products of one reaction priming the products of another reaction.

The term "sexual PCR mutagenesis" or "DNA shuffling" refers to a method of error-prone PCR coupled with forced homologous recombination between DNA molecules of different but highly related DNA sequence in vitro, caused by random fragmentation of the DNA molecule based on sequence similarity, followed by fixation of the crossover by primer extension in an error-prone PCR reaction. See, e.g., Stemmer, Proc. Natl. Acad. Sci. U.S.A. 91: 10747-10751 (1994). DNA shuffling can be carried out between several related genes ("Family shuffling").

The term "in vivo mutagenesis" refers to a process of generating random mutations in any cloned DNA of interest which involves the propagation of the DNA in a strain of bacteria such as E. coli that carries mutations in one or more of the DNA repair pathways. These "mutator" strains have a higher random mutation rate than that of a wild-type parent. Propagating the DNA in a mutator strain will eventually generate random mutations within the DNA.

The term "cassette mutagenesis" refers to any process for replacing a small region of a double-stranded DNA molecule with a synthetic oligonucleotide "cassette" that differs from the native sequence. The oligonucleotide often contains completely and/or partially randomized native sequence.

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The term "recursive ensemble mutagenesis" refers to an algorithm for protein engineering (protein mutagenesis) developed to produce diverse populations of phenotypically related mutants whose members differ in amino acid sequence. This method uses a feedback mechanism to control successive rounds of combinatorial cassette mutagenesis. See, e.g., Arkin et al., Proc. Natl. Acad. Sci. U.S.A. 89: 7811-7815 (1992).

The term "exponential ensemble mutagenesis" refers to a process for generating combinatorial libraries with a high percentage of unique and functional mutants, wherein small groups of residues are randomized in parallel to identify, at each altered position, amino acids which lead to functional proteins. See, e.g., Delegrave et al., Biotechnology Research 11: 1548-1552 (1993); Amold, Current Opinion in Biotechnology 4: 450-455 (1993).

"Operatively linked" expression control sequences refers to a linkage in which the expression control sequence is either contiguous with the gene of interest to control the gene of interest, or acts in *trans* or at a distance to control the gene of interest.

The term "expression control sequence" as used herein refers to polynucleotide sequences which are necessary to affect the expression of coding sequences to which they are operatively linked. Expression control sequences are sequences which control the transcription, post-transcriptional events and translation of nucleic acid sequences. Expression control sequences include appropriate transcription initiation, termination, promoter and enhancer sequences; efficient RNA processing signals such as splicing and polyadenylation signals; sequences that stabilize cytoplasmic mRNA; sequences that enhance translation efficiency (e.g., ribosome binding sites); sequences that enhance protein stability; and when desired, sequences that enhance protein stability; and when desired, sequences that enhance protein stability.

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of such control sequences differs depending upon the host organism; in prokaryotes, such control sequences generally include promoter, ribosomal binding site, and transcription termination sequence. The term "control sequences" is intended to include, at a minimum, all components whose presence is essential for expression, and can also include additional components whose presence is advantageous, for example, leader sequences and fusion partner sequences.

The term "vector," as used herein, is intended to refer to a nucleic acid molecule capable of transporting another nucleic acid to which it has been linked. One type of vector is a "plasmid", which refers to a circular double stranded DNA loop into which additional DNA segments may be ligated. Other vectors include cosmids, bacterial artificial chromosomes (BAC) and yeast artificial chromosomes (YAC). Another type of vector is a viral vector, wherein additional DNA segments may be ligated into the viral genome. Viral vectors that infect bacterial cells are referred to as bacteriophages. Certain vectors are capable of autonomous replication in a host cell into which they are introduced 15 (e.g., bacterial vectors having a bacterial origin of replication). Other vectors can be integrated into the genome of a host cell upon introduction into the host cell, and thereby are replicated along with the host genome. Moreover, certain vectors are canable of directing the expression of genes to which they are operatively linked. Such vectors are referred to herein as "recombinant expression vectors" (or simply, "expression vectors"). In general, expression vectors of utility in recombinant DNA techniques are often in the form of plasmids. In the present specification, "plasmid" and "vector" may be used interchangeably as the plasmid is the most commonly used form of vector. However, the invention is intended to include other forms of expression vectors that serve equivalent functions

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The term "recombinant host cell" (or simply "host cell"), as used herein, is intended to refer to a cell into which a recombinant expression vector has been introduced. It should be understood that such terms are intended to refer not only to the particular subject cell but to the progeny of such a cell. Because certain modifications may occur in succeeding generations due to either mutation or environmental influences, such progeny may not, in fact, be identical to the parent cell, but are still included within the scope of the term "host cell" as used herein.

As used herein, the phrase "open reading frame" and the equivalent acronym "ORF" refers to that portion of a transcript-derived nucleic acid that can be translated in its

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entirety into a sequence of contiguous amino acids. As so defined, an ORF has length, measured in nucleotides, exactly divisible by 3. As so defined, an ORF need not encode the entirety of a natural protein.

As used herein, the phrase "ORF-encoded peptide" refers to the predicted or actual translation of an ORF.

As used herein, the phrase "degenerate variant" of a reference nucleic acid sequenceis meant to be inclusive of all nucleic acid sequences that can be directly translated, using the standard genetic code, to provide an amino acid sequence identical to that translated from the reference nucleic acid sequence.

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The term "polypeptide" encompasses both naturally-occurring and non-naturallyoccurring proteins and polypeptides, as well as polypeptide fragments and polypeptide mutants, derivatives and analogs thereof. A polypeptide may be monomeric or polymeric. Further, a polypeptide may comprise a number of different modules within a single polypeptide each of which has one or more distinct activities. A preferred polypeptide in accordance with the invention comprises a CSP encoded by a nucleic acid molecule of the instant invention, or s a fragment, mutant, analog and derivative thereof.

The term "isolated protein" or "isolated polypeptide" is a protein or polypeptide that by virtue of its origin or source of derivation (1) is not associated with naturally associated components that accompany it in its native state, (2) is free of other proteins from the same species (3) is expressed by a cell from a different species, or (4) does not occur in nature. Thus, a polypeptide that is chemically synthesized or synthesized in a cellular system different from the cell from which it naturally originates will be "isolated" from its naturally associated components. A polypeptide or protein may also be rendered substantially free of naturally associated components by isolation, using protein purification techniques well-known in the art.

A protein or polypeptide is "substantially pure," "substantially homogeneous" or "substantially purified" when at least about 60% to 75% of a sample exhibits a single species of polypeptide. The polypeptide or protein may be monomeric or multimeric. A substantially pure polypeptide or protein will typically comprise about 50%, 60%, 70%, 80% or 90% W/W of a protein sample, more usually about 95%, and preferably will be over 99% pure. Protein purity or homogeneity may be determined by a number of means well-known in the art, such as polyacrylamide gel electrophoresis of a protein sample, followed by visualizing a single polypeptide band upon staining the gel with a stain well-

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known in the art. For certain purposes, higher resolution may be provided by using HPLC or other means well-known in the art for purification.

The term "fragment" when used herein with respect to polypeptides of the present invention invention refers to a polypeptide that has an amino-terminal and/or carboxy-terminal deletion compared to a full-length CSP. In a preferred embodiment, the fragment is a contiguous sequence in which the amino acid sequence of the fragment is identical to the corresponding positions in the naturally-occurring polypeptide. Fragments

typically are at least 5, 6, 7, 8, 9 or 10 amino acids long, preferably at least 12, 14, 16 or 18 amino acids long, more preferably at least 20 amino acids long, more preferably at least 25, 30, 35, 40 or 45, amino acids, even more preferably at least 50 or 60 amino acids long, and even more preferably at least 70 amino acids long.

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A "derivative" when used herein with respect to polypeptidesof the present invention refers to a polypeptide which is substantially similar in primary structural sequence to a CSP but which include, e.g., in vivo or in vitro chemical and biochemical modifications that are not found in the CSP. Such modifications include, for example, acetylation, acylation, ADP-ribosylation, amidation, covalent attachment of flavin, covalent attachment of a heme moiety, covalent attachment of a nucleotide or nucleotide derivative, covalent attachment of a lipid or lipid derivative, covalent attachment of phosphotidylinositol, cross-linking, cyclization, disulfide bond formation, demethylation, formation of covalent cross-links, formation of cystine, formation of pyroglutamate, formylation, gamma-carboxylation, glycosylation, GPI anchor formation, hydroxylation,

iodination, methylation, myristoylation, oxidation, proteolytic processing,
phosphorylation, prenylation, racemization, selenoylation, sulfation, transfer-RNA
mediated addition of amino acids to proteins such as arginylation, and ubiquitination.

25 Other modification include, e.g., labeling with radionuclides, and various enzymatic

modifications, as will be readily appreciated by those skilled in the art. A variety of methods for labeling polypeptides and of substituents or labels useful for such purposes are well-known in the art, and include radioactive isotopes such as ¹²⁵1, ³²P, ³⁵S, ¹⁴C and ³H, ligands which bind to labeled antiligands (e.g., antibodies), fluorophores,

30 chemiluminescent agents, enzymes, and antiligands which can serve as specific binding pair members for a labeled ligand. The choice of label depends on the sensitivity required, ease of conjugation with the primer, stability requirements, and available instrumentation.

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Methods for labeling polypeptides are well-known in the art. See Ausubel (1992), supra; Ausubel (1999), supra.

The term "fusion protein" refers to polypeptides of the present invention coupled to a heterologous amino acid sequences. Fusion proteins are useful because they can be constructed to contain two or more desired functional elements from two or more different proteins. A fusion protein comprises at least 10 contiguous amino acids from a polypeptide of interest, more preferably at least 20 or 30 amino acids, even more preferably at least 40, 50 or 60 amino acids, yet more preferably at least 75, 100 or 125 amino acids. Fusion proteins can be produced recombinantly by constructing a nucleic acid sequence that encodes the polypeptide or a fragment thereof in frame with a nucleic acid sequence encoding a different protein or peptide and then expressing the fusion protein. Alternatively, a fusion protein can be produced chemically by crosslinking the polypeptide or a fragment thereof to another protein.

The term "analog" refers to both polypeptide analogs and non-peptide analogs. The term "polypeptide analog" as used herein refers to a polypeptide that is comprised of a segment of at least 25 amino acids that has substantial identity to a portion of an amino acid sequence but which contains non-natural amino acids or non-natural inter-residue bonds. In a preferred embodiment, the analog has the same or similar biological activity as the native polypeptide. Typically, polypeptide analogs comprise a conservative amino acid substitution (or insertion or deletion) with respect to the naturally-occurring sequence. Analogs typically are at least 20 amino acids long, preferably at least 50 amino acids long or longer, and can often be as long as a full-length naturally-occurring polypeptide.

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The term "non-peptide analog" refers to a compound with properties that are analogous to those of a reference polypeptide. A non-peptide compound may also be termed a "peptide mimetic" or a "peptidomimetic," Such compounds are often developed with the aid of computerized molecular modeling. Peptide mimetics that are structurally similar to useful peptides may be used to produce an equivalent effect. Generally, peptidomimetics are structurally similar to a paradigm polypeptide (i.e., a polypeptide that has a desired biochemical property or pharmacological activity), but have one or more 30 peptide linkages optionally replaced by a linkage selected from the group consisting of: --CH2NH--, --CH2S--, --CH2-CH2--, --CH=CH--(cis and trans), --COCH2--, --CH(OH)CH2--, and -CH2SO--, by methods well-known in the art. Systematic substitution of one or more amino acids of a consensus sequence with a D-amino acid of

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the same type (e.g., D-lysine in place of L-lysine) may also be used to generate more stable peptides. In addition, constrained peptides comprising a consensus sequence or a substantially identical consensus sequence variation may be generated by methods known in the art (Rizo et al., Ann. Rev. Biochem. 61:387-418 (1992)). For example, one may add internal cysteine residues capable of forming intramolecular disulfide bridges which cyclize the pentide.

The term "mutant" or "mutein" when referring to a polypeptide of the present invention relates to an amino acid sequence containing substitutions, insertions or deletions of one or more amino acids compared to the amino acid sequence of a CSP. A mutein may have one or more amino acid point substitutions, in which a single amino acid at a position has been changed to another amino acid, one or more insertions and/or deletions, in which one or more amino acids are inserted or deleted, respectively, in the sequence of the naturally-occurring protein, and/or truncations of the amino acid sequence at either or both the amino or carboxy termini. Further, a mutein may have the same or different biological activity as the naturally-occurring protein. For instance, a mutein may have an increased or decreased biological activity. A mutein has at least 50% sequence similarity to the wild type protein, preferred is 60% sequence similarity, more preferred is 70% sequence similarity. Even more preferred are muteins having 80%, 85% or 90% sequence similarity to a CSP. In an even more preferred embodiment, a mutein exhibits 95% sequence identity, even more preferably 97%, even more preferably 98% and even more preferably 99%. Sequence similarity may be measured by any common sequence analysis algorithm, such as Gap or Bestfit,

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Preferred amino acid substitutions are those which: (1) reduce susceptibility to proteolysis, (2) reduce susceptibility to oxidation, (3) alter binding affinity for forming protein complexes, (4) alter binding affinity or enzymatic activity, and (5) confer or modify other physicochemical or functional properties of such analogs. For example, single or multiple amino acid substitutions (preferably conservative amino acid substitutions) may be made in the naturally-occurring sequence (preferably in the portion of the polypeptide outside the domain(s) forming intermolecular contacts. In a preferred embodiment, the amino acid substitutions are moderately conservative substitutions or conservative substitutions. In a more preferred embodiment, the amino acid substitutions are conservative substitutions. A conservative amino acid substitution should not substantially change the structural characteristics of the parent sequence (e.g., a

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replacement amino acid should not tend to disrupt a helix that occurs in the parent sequence, or disrupt other types of secondary structure that characterizes the parent sequence). Examples of art-recognized polypeptide secondary and tertiary structures are described in Creighton (ed.), Principles, W. H.

Freeman and Company (1984); Branden et al. (ed.), <u>Introduction to Protein Structure</u>, Garland Publishing (1991); Thornton et al., Nature 354:105-106 (1991).

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As used herein, the twenty conventional amino acids and their abbreviations follow conventional usage. See Golub et al. (eds.), Immunology - A Synthesis 2^{nd} Ed., Sinauer Associates (1991). Stereoisomers (e.g., D-amino acids) of the twenty conventional amino acids, unnatural amino acids such as α , α -disubstituted amino acids, N-alkyl amino acids, and other unconventional amino acids may also be suitable components for polypeptides of the present invention. Examples of unconventional amino acids include: 4-hydroxyproline, γ -carboxyglutamate, ε -N,N,N-trimethyllysine, ε -N-acetyllysine, O-phosphoserine, N-acetylserine, N-formylmethionine, 3-methyllistique, of the propagation of

4-hydroxyproline). In the polypeptide notation used herein, the lefthand direction is the amino terminal direction and the right hand direction is the carboxy-terminal direction, in accordance with standard usage and convention.

By "homology" or "homology: when referring to a polypeptide of the present.

By "homology" or "homologous" when referring to a polypeptide of the present invention it is meant polypeptides from different organisms with a similar sequence to the encoded amino acid sequence of a CSP and a similar biological activity or function. Although two polypeptides are said to be "homologous," this does not imply that there is necessarily an evolutionary relationship between the polypeptides. Instead, the term "homologous" is defined to mean that the two polypeptides have similar amino acid sequences and similar biological activities or functions. In a preferred embodiment, a homologous polypeptide is one that exhibits 50% sequence similarity to CSP, preferred is 60% sequence similarity, more preferred is 70% sequence similarity. Even more preferred are homologous polypeptides that exhibit 80%, 85% or 90% sequence similarity to a CSP. In a yet more preferred embodiment, a homologous polypeptide exhibits 95%, 97%, 98% or 99% sequence similarity.

When "sequence similarity" is used in reference to polypeptides, it is recognized that residue positions that are not identical often differ by conservative amino acid substitutions. In a preferred embodiment, a polypeptide that has "sequence similarity"

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comprises conservative or moderately conservative amino acid substitutions. A "conservative amino acid substitution" is one in which an amino acid residue is substituted by another amino acid residue having a side chain (R group) with similar chemical properties (e.g., charge or hydrophobicity). In general, a conservative amino acid substitution will not substantially change the functional properties of a protein. In cases where two or more amino acid sequences differ from each other by conservative substitutions, the percent sequence identity or degree of similarity may be adjusted upwards to correct for the conservative nature of the substitution. Means for making this adjustment are well-known to those of skill in the art. See, e.g., Pearson, Methods Mol. Biol. 24: 307-31 (1994).

For instance, the following six groups each contain amino acids that are conservative substitutions for one another:

- Serine (S), Threonine (T);
- 2) Aspartic Acid (D), Glutamic Acid (E);
- Asparagine (N), Glutamine (Q);

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Arginine (R), Lysine (K);

6.1. Other programs include FASTA, discussed supra.

- 5) Isoleucine (I), Leucine (L), Methionine (M), Alanine (A), Valine (V), and
- Phenylalanine (F), Tyrosine (Y), Tryptophan (W).

Alternatively, a conservative replacement is any change having a positive value in the PAM250 log-likelihood matrix disclosed in Gonnet et al., Science 256: 1443-45 (1992). A "moderately conservative" replacement is any change having a nonnegative value in the PAM250 log-likelihood matrix.

Sequence similarity for polypeptides, which is also referred to as sequence identity, is typically measured using sequence analysis software. Protein analysis software matches similar sequences using measures of similarity assigned to various substitutions, deletions and other modifications, including conservative amino acid substitutions. For instance, GCG contains programs such as "Gap" and "Bestfit" which can be used with default parameters to determine sequence homology or sequence identity between closely related polypeptides, such as homologous polypeptides from different species of organisms or between a wild type protein and a mutein thereof. See, e.g., GCG Version

A preferred algorithm when comparing a sequence of the invention to a database containing a large number of sequences from different organisms is the computer program

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28 BLAST, especially blastp or tblastn. See, e.g., Altschul et al., J. Mol. Biol. 215: 403-410 (1990); Altschul et al., Nucleic Acids Res. 25:3389-402 (1997). Preferred parameters for blastp are:

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Expectation value: 10 (default) 5 Filter: seg (default) Cost to open a gap: 11 (default) Cost to extend a gap: 1 (default Max. alignments: 100 (default) Word size: 11 (default)

No. of descriptions: 100 (default) 10 Penalty Matrix: BLOSUM62

> The length of polypeptide sequences compared for homology will generally be at least about 16 amino acid residues, usually at least about 20 residues, more usually at least about 24 residues, typically at least about 28 residues, and preferably more than about 35 residues. When searching a database containing sequences from a large number of different organisms, it is preferable to compare amino acid sequences.

Algorithms other than blasto for database searching using amino acid sequences are known in the art. For instance, polypeptide sequences can be compared using FASTA, a program in GCG Version 6.1. FASTA (e.g., FASTA2 and FASTA3) provides alignments and percent sequence identity of the regions of the best overlap between the query and search sequences (Pearson (1990), supra; Pearson (2000), supra. For example, percent sequence identity between amino acid sequences can be determined using FASTA with its default or recommended parameters (a word size of 2 and the PAM250 scoring matrix), as provided in GCG Version 6.1.

An "antibody" refers to an intact immunoglobulin, or to an antigen-binding portion thereof that competes with the intact antibody for specific binding to a molecular species, e.g., a polypeptide of the instant invention. Antigen-binding portions may be produced by recombinant DNA techniques or by enzymatic or chemical cleavage of intact antibodies. Antigen-binding portions include, inter alia, Fab, Fab', F(ab')2 Fv, dAb, and complementarity determining region (CDR) fragments, single-chain antibodies (scFv). chimeric antibodies, diabodies and polypeptides that contain at least a portion of an

immunoglobulin that is sufficient to confer specific antigen binding to the polypeptide. A Fab fragment is a monovalent fragment consisting of the VL, VH, CL and CH1 domains;

a F(ab')₂ fragment is a bivalent fragment comprising two Fab fragments linked by a disulfide bridge at the hinge region; a Fd fragment consists of the VH and CH1 domains; a Fv fragment consists of the VL and VH domains of a single arm of an antibody; and a dAb fragment consists of a VH domain, See, e.g., Ward et al., Nature 341: 544-546 (1989).

By "bind specifically" and "specific binding" as used herein it is meant the ability of the antibody to bind to a first molecular species in preference to binding to other molecular species with which the antibody and first molecular species are admixed. An antibody is said specifically to "recognize" a first molecular species when it can bind specifically to that first molecular species.

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A single-chain antibody (scFv) is an antibody in which VL and VH regions are paired to form a monovalent molecule via a synthetic linker that enables them to be made as a single protein chain. See, e.g., Bird et al., Science 242: 423-426 (1988); Huston et al., Proc. Natl. Acad. Sci. USA 85: 5879-5883 (1988). Diabodies are bivalent, bispecific antibodies in which VH and VL domains are expressed on a single polypeptide chain, but using a linker that is too short to allow for pairing between the two domains on the same chain, thereby forcing the domains to pair with complementary domains of another chain and creating two antigen binding sites. See e.g., Holliger et al., Proc. Natl. Acad. Sci. USA 90: 6444-6448 (1993); Poliak et al., Structure 2: 1121-1123 (1994). One or more CDRs may be incorporated into a molecule either covalently or noncovalently to make it an immunoadhesin. An immunoadhesin may incorporate the CDR(s) as part of a larger polypeptide chain, may covalently link the CDR(s) to another polypeptide chain, or may incorporate the CDR(s) noncovalently. The CDRs permit the immunoadhesin to specifically bind to a particular antigen of interest. A chimeric antibody is an antibody that contains one or more regions from one antibody and one or more regions from one or more other antibodies

An antibody may have one or more binding sites. If there is more than one binding site, the binding sites may be identical to one another or may be different. For instance, a naturally-occurring immunoglobulin has two identical binding sites, a single-chain antibody or Fab fragment has one binding site, while a "bispecific" or "bifunctional" antibody has two different binding sites.

An "isolated antibody" is an antibody that (1) is not associated with naturallyassociated components, including other naturally-associated antibodies, that accompany it in its native state, (2) is free of other proteins from the same species, (3) is expressed by a

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cell from a different species, or (4) does not occur in nature. It is known that purified proteins, including purified antibodies, may be stabilized with non-naturally-associated components. The non-naturally-associated component may be a protein, such as albumin (e.g., BSA) or a chemical such as polyethylene glycol (PEG).

A "neutralizing antibody" or "an inhibitory antibody" is an antibody that inhibits the activity of a polypeptide or blocks the binding of a polypeptide to a ligand that normally binds to it. An "activating antibody" is an antibody that increases the activity of a polypeptide.

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The term "epitope" includes any protein determinant capable of specific binding to 10 an immunoglobulin or T-cell receptor. Epitopic determinants usually consist of chemically active surface groupings of molecules such as amino acids or sugar side chains and usually have specific three dimensional structural characteristics, as well as specific charge characteristics. An antibody is said to specifically bind an antigen when the dissociation constant is less than 1 µM, preferably less than 100 nM and most preferably less than 10 nM.

The term "patient" includes human and veterinary subjects.

Throughout this specification and claims, the word "comprise," or variations such as "comprises" or "comprising," will be understood to imply the inclusion of a stated integer or group of integers but not the exclusion of any other integer or group of integers.

The term "colon specific" refers to a nucleic acid molecule or polypentide that is expressed predominantly in the colon as compared to other tissues in the body. In a preferred embodiment, a "colon specific" nucleic acid molecule or polypeptide is detected at a level that is 2-fold higher than any other tissue in the body. In a more preferred embodiment, the "colon specific" nucleic acid molecule or polypeptide is detected at a level that is 5-fold higher than any other tissue in the body, more preferably at least 10fold, 15-fold, 20-fold, 25-fold, 50-fold or 100-fold higher than any other tissue in the body. Nucleic acid molecule levels may be measured by nucleic acid hybridization, such as Northern blot hybridization, or quantitative PCR. Polypeptide levels may be measured by any method known to accurately quantitate protein levels, such as Western blot analysis.

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Nucleic Acid Molecules, Regulatory Sequences, Vectors, Host Cells and Recombinant Methods of Making Polypeptides

Nucleic Acid Molecules

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One aspect of the invention provides isolated nucleic acid molecules that are specific to the colon or to colon cells or tissue or that are derived from such nucleic acid molecules. These isolated colon specific nucleic acids (CSNAs) may comprise cDNA genomic DNA, RNA, or a combination thereof, a fragment of one of these nucleic acids, or may be a non-naturally-occurring nucleic acid molecule. A CSNA may be derived from an animal. In a preferred embodiment, the CSNA is derived from a human or other 10 mammal. In a more preferred embodiment, the CSNA is derived from a human or other primate. In an even more preferred embodiment, the CSNA is derived from a human.

In a preferred embodiment, the nucleic acid molecule encodes a polypeptide that is specific to colon, a colon-specific polypeptide (CSP). In a more preferred embodiment, the nucleic acid molecule encodes a polypeptide that comprises an amino acid sequence of SEQ ID NO: 101-190. In another highly preferred embodiment, the nucleic acid molecule comprises a nucleic acid sequence of SEQ ID NO: 1-100. Nucleotide sequences of the instantly-described nucleic acid molecules were determined by sequencing a DNA molecule that had resulted, directly or indirectly, from at least one enzymatic polymerization reaction (e.g., reverse transcription and/or polymerase chain reaction) using an automated sequencer (such as the MegaBACETM 1000, Amersham Biosciences, Sunnyvale, CA, USA).

Nucleic acid molecules of the present invention may also comprise sequences that selectively hybridizes to a nucleic acid molecule encoding a CSNA or a complement or antisense thereof. The hybridizing nucleic acid molecule may or may not encode a polypeptide or may or may not encode a CSP. However, in a preferred embodiment, the hybridizing nucleic acid molecule encodes a CSP. In a more preferred embodiment, the invention provides a nucleic acid molecule that selectively hybridizes to a nucleic acid molecule or the antisense sequence of a nucleic acid molecule that encodes a polypeptide comprising an amino acid sequence of SEO ID NO: 101-190. In an even more preferred embodiment, the invention provides a nucleic acid molecule that selectively hybridizes to a nucleic acid molecule comprising the nucleic acid sequence of SEQ ID NO: 1-100 or the antisense sequence thereof. Preferably, the nucleic acid molecule selectively hybridizes to a nucleic acid molecule or the antisense sequence of a nucleic acid molecule encoding a

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CSP under low stringency conditions. More preferably, the nucleic acid molecule selectively hybridizes to a nucleic acid molecule or the antisense sequence of a nucleic acid molecule encoding a CSP under moderate stringency conditions. Most preferably, the nucleic acid molecule selectively hybridizes to a nucleic acid molecule or the antisense sequence of a nucleic acid molecule encoding a CSP under high stringency conditions. In a preferred embodiment, the nucleic acid molecule hybridizes under low, moderate or high stringency conditions to a nucleic acid molecule or the antisense sequence of a nucleic acid molecule encoding a polypeptide comprising an amino acid sequence of SEQ ID NO: 101-190. In a more preferred embodiment, the nucleic acid molecule hybridizes under low, moderate or high stringency conditions to a nucleic acid molecule or the antisense sequence of a nucleic acid molecule comprising a nucleic acid sequence selected from SEO ID NO: 1-100.

Nucleic acid molecules of the present invention may also comprise nucleic acid sequences that exhibit substantial sequence similarity to a nucleic acid encoding a CSP or a complement of the encoding nucleic acid molecule. In this embodiment, it is preferred that the nucleic acid molecule exhibit substantial sequence similarity to a nucleic acid molecule encoding human CSP. More preferred is a nucleic acid molecule exhibiting substantial sequence similarity to a nucleic acid molecule encoding a polypeptide having an amino acid sequence of SEQ ID NO: 101-190. By substantial sequence similarity it is meant a nucleic acid molecule having at least 60% sequence identity with a nucleic acid molecule encoding a CSP, such as a polypeptide having an amino acid sequence of SEO ID NO: 101-190, more preferably at least 70%, even more preferably at least 80% and even more preferably at least 85%. In a more preferred embodiment, the similar nucleic acid molecule is one that has at least 90% sequence identity with a nucleic acid molecule encoding a CSP, more preferably at least 95%, more preferably at least 97%, even more preferably at least 98%, and still more preferably at least 99%. Most preferred in this embodiment is a nucleic acid molecule that has at least 99.5%, 99.6%, 99.7%, 99.8% or 99.9% sequence identity with a nucleic acid molecule encoding a CSP.

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The nucleic acid molecules of the present invention are also inclusive of those exhibiting substantial sequence similarity to a CSNA or its complement. In this embodiment, it is preferred that the nucleic acid molecule exhibit substantial sequence similarity to a nucleic acid molecule having a nucleic acid sequence of SEQ ID NO: 1-100. By substantial sequence similarity it is meant a nucleic acid molecule that has at

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least 60% sequence identity with a CSNA, such as one having a nucleic acid sequence of SEQ ID NO: 1-100, more preferably at least 70%, even more preferably at least 80% and even more preferably at least 85%. More preferred is a nucleic acid molecule that has at least 90% sequence identity with a CSNA, more preferably at least 95%, more preferably at least 99%, even more preferably at least 99%. Most preferred is a nucleic acid molecule that has at least 99.5%, 99.6%, 99.7%, 99.8% or 99.9% sequence identity with a CSNA.

Nucleic acid molecules that exhibit substantial sequence similarity are inclusive of sequences that exhibits sequence identity over their entire length to a CSNA or to a nucleic acid molecule encoding a CSP, as well as sequences that are similar over only a part of its length. In this case, the part is at least 50 nucleotides of the CSNA or the nucleic acid molecule encoding a CSP, preferably at least 100 nucleotides, more preferably at least 150 or 200 nucleotides, even more preferably at least 250 or 300 nucleotides, still more preferably at least 400 or 500 nucleotides.

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15 The substantially similar nucleic acid molecule may be a naturally-occurring one that is derived from another species, especially one derived from another primate, wherein the similar nucleic acid molecule encodes an amino acid sequence that exhibits significant sequence identity to that of SEQ ID NO: 101-190 or demonstrates significant sequence identity to the nucleotide sequence of SEO ID NO: 1-100. The similar nucleic acid molecule may also be a naturally-occurring nucleic acid molecule from a human, when the CSNA is a member of a gene family. The similar nucleic acid molecule may also be a naturally-occurring nucleic acid molecule derived from a non-primate, mammalian species, including without limitation, domesticated species, e.g., dog, cat, mouse, rat, rabbit, hamster, cow, horse and pig; and wild animals, e.g., monkey, fox, lions, tigers, 2.5 bears, giraffes, zebras, etc. The substantially similar nucleic acid molecule may also be a naturally-occurring nucleic acid molecule derived from a non-mammalian species, such as birds or reptiles. The naturally-occurring substantially similar nucleic acid molecule may be isolated directly from humans or other species. In another embodiment, the substantially similar nucleic acid molecule may be one that is experimentally produced by 30 random mutation of a nucleic acid molecule. In another embodiment, the substantially similar nucleic acid molecule may be one that is experimentally produced by directed mutation of a CSNA. In a preferred embodiment, the substantially similar nucleic acid molecule is an CSNA.

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The nucleic acid molecules of the present invention are also inclusive of allelic variants of a CSNA or a nucleic acid encoding a CSP. For example, single nucleotide polymorphisms (SNPs) occur frequently in eukaryotic genomes and the sequence determined from one individual of a species may differ from other allelic forms present within the population. More than 1.4 million SNPs have already identified in the human genome, International Human Genome Sequencing Consortium, Nature 409: 860-921 (2001) — Variants with small deletions and insertions of more than a single nucleotide are also found in the general population, and often do not alter the function of the protein. In addition, amino acid substitutions occur frequently among natural allelic variants, and often do not substantially change protein function.

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In a preferred embodiment, the allelic variant is a variant of a gene, wherein the gene is transcribed into an mRNA that encodes a CSP. In a more preferred embodiment, the gene is transcribed into an mRNA that encodes a CSP comprising an amino acid sequence of SEQ ID NO: 101-190. In another preferred embodiment, the allelic variant is a variant of a gene, wherein the gene is transcribed into an mRNA that is a CSNA. In a more preferred embodiment, the gene is transcribed into an mRNA that comprises the nucleic acid sequence of SEQ ID NO: 1-100. Also preferred is that the allelic variant is a naturally-occurring allelic variant in the species of interest, particularly human.

Nucleic acid molecules of the present invention are also inclusive of nucleic acid sequences comprising a part of a nucleic acid sequence of the instant invention. The part may or may not encode a polypeptide, and may or may not encode a polypeptide that is a CSP. In a preferred embodiment, the part encodes a CSP. In one embodiment, the nucleic acid molecule comprises a part of a CSNA. In another embodiment, the nucleic acid molecule comprises a part of a nucleic acid molecule that hybridizes or exhibits substantial sequence similarity to a CSNA. In another embodiment, the nucleic acid molecule comprises a part of a nucleic acid molecule that is an allelic variant of a CSNA. In yet another embodiment, the nucleic acid molecule that is an allelic variant of a CSNA. In yet another embodiment, the nucleic acid molecule comprises a part of a nucleic acid molecule that encodes a CSP. A part comprises at least 10 nucleotides, more preferably at least 15, 17, 18, 20, 25, 30, 35, 40, 50, 60, 70, 80, 90, 100, 150, 200, 250, 300, 350, 400 or 500 nucleotides. The maximum size of a nucleic acid part is one nucleotide shorter than the sequence of the nucleic acid molecule encoding the full-length protein.

Nucleic acid molecules of the present invention are also inclusive of nucleic acid sequences that encode fusion proteins, homologous proteins, polypeptide fragments, muteins and polypeptide analogs, as described infra.

Nucleic acid molecules of the present invention are also inclusive of nucleic acid sequences containing modifications of the native nucleic acid molecule. Examples of such modifications include, but are not limited to, nonnative internucleoside bonds, post-synthetic modifications or altered nucleotide analogues. One having ordinary skill in the art would recognize that the type of modification that can be made will depend upon the intended use of the nucleic acid molecule. For instance, when the nucleic acid molecule is used as a hybridization probe, the range of such modifications will be limited to those that permit sequence-discriminating base pairing of the resulting nucleic acid. When used to direct expression of RNA or protein in vitro or in vivo, the range of such modifications will be limited to those that permit the nucleic acid to function properly as a polymerization substrate. When the isolated nucleic acid is used as a therapeutic agent, the modifications will be limited to those that do not confer toxicity upon the isolated nucleic acid.

Accordingly, in one embodiment, a nucleic acid molecule may include nucleotide analogues that incorporate labels that are directly detectable, such as radiolabels or fluorophores, or nucleotide analogues that incorporate labels that can be visualized in a subsequent reaction, such as biotin or various haptens. The labeled nucleic acid molecules are particularly useful as hybridization probes.

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Common radiolabeled analogues include those labeled with ³³P, ³²P, and ³⁵S, such as α-²²P-dATP, α-³²P-dCTP, α-³²P-dGTP, α-³²P-dTTP, α-³²P-3³dATP, α-³²P-ATP, α-³²P-CTP, α-²²P-GTP, α-²²P-UTP, α-²⁵S-GTP, γ-²⁵S-GTP, γ-²³P-dATP, and the like.

Commercially available fluorescent nucleotide analogues readily incorporated into the nucleic acids of the present invention include Cy3-dCTP, Cy3-dUTP, Cy5-dUTP (Amersham Biosciences, Piscataway, New Jersey, USA), fluorescein-12-dUTP, tetramethylrhodamine-6-dUTP, Texas Red®-5-dUTP, Cascade Blue®-7-dUTP, BODIPY® FL-14-dUTP, BODIPY® FL-14-dUTP, BODIPY® TR-14-dUTP.

30 Rhodamine Green™-5-dUTP, Oregon Green® 488-5-dUTP, Texas Red®-12-dUTP, BODIPY® 630/650-14-dUTP, BODIPY® 650/665-14-dUTP, Alexa Fluor® 488-5-dUTP, Alexa Fluor® 532-5-dUTP, Alexa Fluor® 568-5-dUTP, Alexa Fluor® 594-5-dUTP, Alexa Fluor® 546-14-dUTP, fluorescein-12-UTP, tetramethylrhodamine-6-UTP, Texas

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Red®-5-UTP, Cascade Blue®-7-UTP, BODIPY® FL-14-UTP, BODIPY® TMR-14-UTP, BODIPY® TR-14-UTP, Rhodamine Green™-5-UTP, Alexa Fluor® 546-14-UTP (Molecular Probes, Inc. Eugene, OR, USA). One may also custom synthesize nucleotides having other fluorophores. See Henegariu et al., Nature Biotechnol. 18: 345-348 (2000).

Haptens that are commonly conjugated to nucleotides for subsequent labeling include biotin (biotin-11-dUTP, Molecular Probes, Inc., Eugene, OR, USA; biotin-21-UTP, biotin-21-dUTP, Clontech Laboratories, Inc., Palo Alto, CA, USA), digoxigenin (DIG-11-dUTP, alkali labile, DIG-11-UTP, Roche Diagnostics Corp., Indianapolis, IN, USA), and dinitrophenyl (dinitrophenyl-11-dUTP, Molecular Probes, Inc., Eugene, OR, USA).

Nucleic acid molecules of the present invention can be labeled by incorporation of labeled nucleotide analogues into the nucleic acid. Such analogues can be incorporated by enzymatic polymerization, such as by nick translation, random priming, polymerase chain reaction (PCR), terminal transferase tailing, and end-filling of overhangs, for DNA molecules, and in vitro transcription driven, e.g., from phage promoters, such as T7, T3, and SP6, for RNA molecules. Commercial kits are readily available for each such labeling approach. Analogues can also be incorporated during automated solid phase chemical synthesis. Labels can also be incorporated after nucleic acid synthesis, with the 5' phosphate and 3' hydroxyl providing convenient sites for post-synthetic covalent attachment of detectable labels.

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Other post-synthetic approaches also permit internal labeling of nucleic acids. For example, fluorophores can be attached using a cisplatin reagent that reacts with the N7 of guanine residues (and, to a lesser extent, adenine bases) in DNA, RNA, and Peptide

25 Nucleic Acids (PNA) to provide a stable coordination complex between the nucleic acid and fluorophore label (Universal Linkage System) (available from Molecular Probes, Inc., Eugene, OR, USA and Amersham Pharmacia Biotech, Piscataway, NJ, USA); see Alers et al., Genes, Chromosomes & Cancer 25: 301-305 (1999); Jelsma et al., J. NIH Res. 5: 82 (1994); Van Belkum et al., BioTechniques 16: 148-153 (1994). Alternatively, nucleic acid scan be labeled using a disulfide-containing linker (FastTag™ Reagent, Vector Laboratories, Inc., Burlingame, CA, USA) that is photo- or thermally coupled to the target nucleic acid using aryl azide chemistry; after reduction, a free thiol is available for coupling to a hapten, fluorophore, sugar, affinity licand, or other marker.

herein by reference in their entireties.

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One or more independent or interacting labels can be incorporated into the nucleic acid molecules of the present invention. For example, both a fluorophore and a moiety that in proximity thereto acts to quench fluorescence can be included to report specific hybridization through release of fluorescence quenching or to report exonucleotidic excision. See, e.g., Tyagi et al., Nature Biotechnol. 14: 303-308 (1996); Tyagi et al., Nature Biotechnol. 16: 49-53 (1998); Sokol et al., Proc. Natl. Acad. Sci. USA 95: 11538-11543 (1998); Kostrikis et al., Science 279: 1228-1229 (1998); Marras et al., Genet. Anal. 14: 151-156 (1999); Holland et al., Proc. Natl. Acad. Sci. USA 88: 7276-7280 (1991); Heid et al., Genome Res. 6(10): 986-94 (1996); Kuimelis et al., Nucleic Acids Symp. Ser. (37): 255-6 (1997); and United States Patent Nos. 5,846,726, 5,925,517, 5,925,517, 5,723,591 and 5,538,848, the disclosures of which are incorporated

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Nucleic acid molecules of the present invention may also be modified by altering one or more native phosphodiester internucleoside bonds to more nuclease-resistant, internucleoside bonds. See Hartmann et al. (eds.), Manual of Antisense Methodology; Perspectives in Antisense Science, Kluwer Law International (1999); Stein et al. (eds.), Applied Antisense Oligonucleotide Technology, Wiley-Liss (1998); Chadwick et al. (eds.), Oligonucleotides as Therapeutic Agents - Symposium No. 209. John Wiley & Son Ltd (1997). Such altered internucleoside bonds are often desired for antisense techniques or for targeted gene correction, Gamper et al., Nucl. Acids Res. 28(21): 4332-4339 (2000).

Modified oligonucleotide backbones include, without limitation, phosphorothioates, chiral phosphorothioates, phosphorodithioates, phosphoriesters, aminoalkylphosphotriesters, methyl and other alkyl phosphonates including 3'-alkylene phosphonates and chiral phosphonates, phosphinates, phosphoramidates including 3'-amino phosphoramidate and aminoalkylphosphoramidates, thionoalkylphosphoramidates, thionoalkylphosphorames, thionoalkylphosphoramidates, and boranophosphates having normal 3'-5' linkages, 2'-5' linked analogs of these, and those having inverted polarity wherein the adjacent pairs of nucleoside units are linked 3'-5' to 5'-3' or 2'-5' to 5'-2'. Representative United States patents that teach the preparation of the above phosphorus-containing linkages include, but are not limited to, United States Patent Nos. 3,687,808; 4,469,863; 4,476,301; 5,023,243; 5,177,196; 5,188,897; 5,264,423; 5,276,019;

5,278,302; 5,286,717; 5,321,131; 5,399,676; 5,405,939; 5,453,496; 5,455,233; 5,466,677; 5,476,925; 5,519,126; 5,536,821; 5,541,306; 5,550,111; 5,563,253; 5,571,799; 5,587,361;

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and 5,625,050, the disclosures of which are incorporated herein by reference in their entireties. In a preferred embodiment, the modified internucleoside linkages may be used for antisense techniques.

Other modified oligonucleotide backbones do not include a phosphorus atom, but have backbones that are formed by short chain alkyl or cycloalkyl internucleoside linkages, mixed heteroatom and alkyl or cycloalkyl internucleoside linkages, or one or more short chain heteroatomic or heterocyclic internucleoside linkages. These include those having morpholino linkages (formed in part from the sugar portion of a nucleoside); siloxane backbones; sulfide, sulfoxide and sulfone backbones; formacetyl and thioformacetyl backbones; methylene formacetyl and thioformacetyl backbones; alkene containing backbones; sulfamate backbones; methyleneimino and methylenehydrazino backbones; sulfonate and sulfonamide backbones; amide backbones; and others having mixed N, O, S and CH₂ component parts. Representative U.S. patents that teach the preparation of the above backbones include, but are not limited to, United States Patent Nos. 5,034,506; 5,166,315; 5,185,444; 5,214,134; 5,216,141; 5,235,033; 5,264,562; 5,264,564; 5,405,938; 5,434,257; 5,466,677; 5,470,967; 5,489,677; 5,541,307; 5,561,225; 5,596,086; 5,602,240; 5,610,289; 5,602,240; 5,602,240; 5,602,290; 5,618,704; 5,623,070;

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5,264,304; 3,403,948; 3,444,257; 3,406,677; 3,470,967; 3,489,677; 3,541,307; 5,561,425; 5,596,086; 5,602,240; 5,610,289; 5,602,240; 5,608,046; 5,601,289; 5,618,704; 5,623,070; 5,663,312; 5,633,360; 5,677,437 and 5,677,439; the disclosures of which are incorporated herein by reference in their entireties.

In other preferrednucleic acid molecules, both the sugar and the internucleoside

linkage are replaced with novel groups, such as peptide nucleic acids (PNA). In PNA compounds, the phosphodiester backbone of the nucleic acid is replaced with an amide-containing backbone, in particular by repeating N-(2-aminoethyl) glycine units linked by amide bonds. Nucleobases are bound directly or indirectly to aza nitrogen atoms of the amide portion of the backbone, typically by methylene carbonyl linkages. PNA can be synthesized using a modified peptide synthesis protocol. PNA oligomers can be synthesized by both Fmoc and tBoc methods. Representative U.S. patents that teach the preparation of PNA compounds include, but are not limited to, United States Patent Nos. 5,539,082; 5,714,331; and 5,719,262, each of which is herein incorporated by reference in its entirety. Automated PNA synthesis is readily achievable on commercial synthesizers (see, e.g., "PNA User's Guide," Rev. 2, February 1998, Perseptive Biosystems Part No. 60138, Applied Biosystems, Inc., Foster City, CA).PNA molecules are advantageous for a number of reasons. First, because the PNA backbone is uncharged, PNA/DNA and

PNA/RNA duplexes have a higher thermal stability than is found in DNA/DNA and DNA/RNA duplexes. The Tm of a PNA/DNA or PNA/RNA duplex is generally 1°C higher per base pair than the Tm of the corresponding DNA/DNA or DNA/RNA duplex (in 100 mM NaCl). Second, PNA molecules can also form stable PNA/DNA complexes at low ionic strength, under conditions in which DNA/DNA duplex formation does not occur. Third, PNA also demonstrates greater specificity in binding to complementary DNA because a PNA/DNA mismatch is more destabilizing than DNA/DNA mismatch. A single mismatch in mixed a PNA/DNA 15-mer lowers the Tm by 8-20°C (15°C on average). In the corresponding DNA/DNA duplexes, a single mismatch lowers the Tm by 10 4-16°C (11°C on average). Because PNA probes can be significantly shorter than DNA probes, their specificity is greater. Fourth, PNA oligomers are resistant to degradation by enzymes, and the lifetime of these compounds is extended both in vivo and in vitro because nucleases and proteases do not recognize the PNA polyamide backbone with nucleobase sidechains. See, e.g., Ray et al., FASEB J. 14(9): 1041-60 (2000); Nielsen et 15 al., Pharmacol Toxicol, 86(1): 3-7 (2000); Larsen et al., Biochim Biophys Acta, 1489(1): 159-66 (1999); Nielsen, Curr. Opin. Struct. Biol. 9(3): 353-7 (1999), and Nielsen, Curr. Opin. Biotechnol. 10(1): 71-5 (1999).

Nucleic acid molecules may be modified compared to their native structure throughout the length of the nucleic acid molecule or can be localized to discrete portions thereof. As an example of the latter, chimeric nucleic acids can be synthesized that have discrete DNA and RNA domains and that can be used for targeted gene repair and modified PCR reactions, as further described in, Misra et al., Biochem. 37: 1917-1925 (1998); and Finn et al., Nucl. Acids Res. 24: 3357-3363 (1996), and United States Patent Nos. 5,760,012 and 5,731,181, the disclosures of which are incorporated herein by reference in their entireties.

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Unless otherwise specified, nucleic acid molecules of the present invention can include any topological conformation appropriate to the desired use; the term thus explicitly comprehends, among others, single-stranded, double-stranded, triplexed, quadruplexed, partially double-stranded, partially-triplexed, partially-quadruplexed, branched, hairpinned, circular, and padlocked conformations. Padlock conformations and their utilities are further described in Banér et al., Curr. Opin. Biotechnol. 12: 11-15 (2001); Escude et al., Proc. Natl. Acad. Sci. USA 14: 96(19):10603-7 (1999); and Nilsson et al., Science 265(5181): 2085-8 (1994). Triplex and quadruplex conformations, and

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their utilities, are reviewed in Praseuth et al., Biochim. Biophys. Acta. 1489(1): 181-206 (1999); Fox, Curr. Med. Chem. 7(1): 17-37 (2000); Kochetkova et al., Methods Mol. Biol. 130: 189-201 (2000); Chan et al., J. Mol. Med. 75(4): 267-82 (1997).

Methods for Using Nucleic Acid Molecules as Probes and Primers

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The isolated nucleic acid molecules of the present invention can be used as hybridization probes to detect, characterize, and quantify hybridizing nucleic acids in, and isolate hybridizing nucleic acids from, both genomic and transcript-derived nucleic acid samples. When free in solution, such probes are typically, but not invariably, detectably labeled; bound to a substrate, as in a microarray, such probes are typically, but not invariably unlabeled.

In one embodiment, the isolated nucleic acid molecules of the present invention can be used as probes to detect and characterize gross alterations in the gene of a CSNA, such as deletions, insertions, translocations, and duplications of the CSNA genomic locus through fluorescence in stu hybridization (FISH) to chromosome spreads. See, e.g., Andreeff et al. (eds.), Introduction to Fluorescence In Stu Hybridization: Principles and Clinical Applications, John Wiley & Sons (1999). The isolated nucleic acid molecules of the present invention can be used as probes to assess smaller genomic alterations using, e.g., Southern blot detection of restriction fragment length polymorphisms. The isolated nucleic acid molecules of the present invention can be used as probes to isolate genomic clones that include a nucleic acid molecule of the present invention, which thereafter can be restriction mapped and sequenced to identify deletions, insertions, translocations, and substitutions (single nucleotide polymorphisms, SNPs) at the sequence level.

The isolated nucleic acid molecules of the present invention can be also be used as probes to detect, characterize, and quantify CSNA in, and isolate CSNA from, transcript-derived nucleic acid samples. In one embodiment, the isolated nucleic acid molecules of the present invention can be used as hybridization probes to detect, characterize by length, and quantify mRNA by Northern blot of total or poly-A*- selected RNA samples. In another embodiment, the isolated nucleic acid molecules of the present invention can be used as hybridization probes to detect, characterize by location, and quantify mRNA by in situ hybridization to tissue sections. See, e.g., Schwarchzacher et al., In Situ Hybridization, Springer-Verlag New York (2000). In another preferred embodiment, the isolated nucleic acid molecules of the present invention can be used as hybridization

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probes to measure the representation of clones in a cDNA library or to isolate hybridizing nucleic acid molecules acids from cDNA libraries, permitting sequence level characterization of mRNAs that hybridize to CSNAs, including, without limitations, identification of deletions, insertions, substitutions, truncations, alternatively spliced forms and single nucleotide polymorphisms. In yet another preferred embodiment, the nucleic acid molecules of the instant invention may be used in microarrays.

All of the aforementioned probe techniques are well within the skill in the art, and are described at greater length in standard texts such as Sambrook (2001), supra; Ausubel (1999), supra; and Walker et al. (eds.), The Nucleic Acids Protocols Handbook, Humana Press (2000).

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In another embodiment, a nucleic acid molecule of the invention may be used as a probe or primer to identify and/or amplify a second nucleic acid molecule that selectively hybridizes to the nucleic acid molecule of the invention. In this embodiment, it is preferred that the probe or primer be derived from a nucleic acid molecule encoding a 15 CSP. More preferably, the probe or primer is derived from a nucleic acid molecule encoding a polypeptide having an amino acid sequence of SEQ ID NO: 101-190. Also preferred are probes or primers derived from a CSNA. More preferred are probes or primers derived from a nucleic acid molecule having a nucleotide sequence of SEQ ID NO: 1100

In general, a probe or primer is at least 10 nucleotides in length, more preferably at least 12, more preferably at least 14 and even more preferably at least 16 or 17 nucleotides in length. In an even more preferad embodiment, the probe or primer is at least 18 nucleotides in length, even more preferably at least 20 nucleotides and even more preferably at least 22 nucleotides in length, even more preferably at least 22 nucleotides in length, or may be 30, 40 or 50 nucleotides in length. Methods of performing nucleic acid hybridization using oligonucleotide probes are well-known in the art. See, e.g., Sambrook et al., 1989, supra, Chapter 11 and pp. 11.31-11.32 and 11.40-11.44, which describes radiolabeling of short probes, and pp. 11.45-11.53, which describe hybridization conditions for oligonucleotide probes, including specific conditions for probe hybridization (pp. 11.50-11.51).

Methods of performing primer-directed amplification are also well-known in the art. Methods for performing the polymerase chain reaction (PCR) are compiled, *inter alia*, in McPherson, PCR Basics: From Background to Bench, Springer Verlag (2000); Innis et

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al. (eds.), PCR Applications: Protocols for Functional Genomics, Academic Press (1999); Gelfand et al. (eds.), PCR Strategies, Academic Press (1998); Newton et al., PCR, Springer-Verlag New York (1997); Burke (ed.), PCR: Essential Techniques, John Wilcy & Son Ltd (1996); White (ed.), PCR Cloning Protocols: From Molecular Cloning to Genetic Engineering, Vol. 67, Humana Press (1996); and McPherson et al. (eds.), PCR 2: A Practical Approach, Oxford University Press, Inc. (1995). Methods for performing RT-PCR are collected, e.g., in Siebert et al. (eds.), Gene Cloning and Analysis by RT-PCR, Eaton Publishing Company/Bio Techniques Books Division, 1998; and Siebert (ed.), PCR Technique: RT-PCR, Eaton Publishing Company/ BioTechniques Books (1995).

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PCR and hybridization methods may be used to identify and/or isolate nucleic acid molecules of the present invention including allelic variants, homologous nucleic acid molecules and fragments. PCR and hybridization methods may also be used to identify. amplify and/or isolate nucleic acid molecules of the present invention that encode homologous proteins, analogs, fusion protein or muteins of the invention. Nucleic acid primers as described herein can be used to prime amplification of nucleic acid molecules of the invention, using transcript-derived or genomic DNA as template.

These nucleic acid primers can also be used, for example, to prime single hase extension (SBE) for SNP detection (See, e.g., U.S. Pat. No. 6,004,744, the disclosure of which is incorporated herein by reference in its entirety).

Isothermal amplification approaches, such as rolling circle amplification, are also now well-described. See, e.g., Schweitzer et al., Curr. Opin. Biotechnol. 12(1): 21-7 (2001); international patent publications WO 97/19193 and WO 00/15779, and United States Patent Nos. 5.854,033 and 5.714,320, the disclosures of which are incorporated herein by reference in their entireties. Rolling circle amplification can be combined with 25 other techniques to facilitate SNP detection. See, e.g., Lizardi et al., Nature Genet. 19(3): 225-32 (1998).

Nucleic acid molecules of the present invention may be bound to a substrate either covalently or noncovalently. The substrate can be porous or solid, planar or non-planar, unitary or distributed. The bound nucleic acid molecules may be used as hybridization probes, and may be labeled or unlabeled. In a preferred embodiment, the bound nucleic acid molecules are unlabeled.

In one embodiment, the nucleic acid molecule of the present invention is bound to a porous substrate, e.g., a membrane, typically comprising nitrocellulose, nylon, or

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positively-charged derivatized nylon. The nucleic acid molecule of the present invention can be used to detect a hybridizing nucleic acid molecule that is present within a labeled nucleic acid sample, e.g., a sample of transcript-derived nucleic acids. In another embodiment, the nucleic acid molecule is bound to a solid substrate, including, without limitation, glass, amorphous silicon, crystalline silicon or plastics. Examples of plastics include, without limitation, polymethylacrylic, polyethylene, polypropylene, polyacrylate, polymethylmethacrylate, polyvinylchloride, polytetrafluoroethylene, polystyrene, polycarbonate, polyacetal, polysulfone, celluloseacetate, cellulosenitrate, nitrocellulose, or mixtures thereof. The solid substrate may be any shape, including rectangular, disk-like and spherical. In a preferred embodiment, the solid substrate is a microscope slide or slide-shaped substrate.

The nucleic acid molecule of the present invention can be attached covalently to a surface of the support substrate or applied to a derivatized surface in a chaotropic agent that facilitates denaturation and adherence by presumed noncovalent interactions, or some combination thereof. The nucleic acid molecule of the present invention can be bound to a substrate to which a plurality of other nucleic acids are concurrently bound, hybridization to each of the plurality of bound nucleic acids being separately detectable. At low density, e.g. on a porous membrane, these substrate-bound collections are typically denominated macroarrays; at higher density, typically on a solid support, such as glass, these substrate bound collections of plural nucleic acids are colloquially termed microarrays. As used herein, the term microarray includes arrays of all densities. It is, therefore, another aspect of the invention to provide microarrays that comprise one or more of the nucleic acid molecules of the present invention.

In yet another embodiment, the invention is directed to single exon probes based on the CSNAs disclosed herein.

Expression Vectors, Host Cells and Recombinant Methods of Producing Polypeptides

Another aspect of the present invention provides vectors that comprise one or more of the isolated nucleic acid molecules of the present invention, and host cells in which such vectors have been introduced.

The vectors can be used, *inter alia*, for propagating the nucleic acid molecules of the present invention in host cells (cloning vectors), for shuttling the nucleic acid

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molecules of the present invention between host cells derived from disparate organisms (shuttle vectors), for inserting the nucleic acid molecules of the present invention into host cell chromosomes (insertion vectors), for expressing sense or antisense RNA transcripts of the nucleic acid moleculess of the present invention in vitro or within a host cell, and for expressing polypeptides encoded by the nucleic acid moleculess of the present invention, alone or as fusion proteins with heterologous polypeptides (expression vectors). Vectors are by now well-known in the art, and are described, inter alia, in Jones et al. (eds.), Vectors: Cloning Applications: Essential Techniques (Essential Techniques Series), John Wiley & Son Ltd. (1998); Jones et al. (eds.), Vectors: Expression Systems: Essential Techniques (Essential Techniques Series), John Wiley & Son Ltd. (1998); Gacesa et al., Vectors: Essential Data, John Wiley & Sons Ltd. (1995); Cid-Arregui (eds.), Viral Vectors: Essential Data, John Wiley & Sons Ltd. (1995); Cid-Arregui (eds.), Viral Vectors: Essential Data, John Wiley & Sons Ltd. (1995); Cid-Arregui (eds.), Viral Vectors: Essential Data, John Wiley & Sons Ltd. (1995); Cid-Arregui (eds.), Viral Vectors: Essential Data, John Wiley & Sons Ltd. (1995); Cid-Arregui (eds.), Viral Vectors: Essential Data, John Wiley & Sons Ltd. (1995); Cid-Arregui (eds.), Viral Vectors: Essential Data, John Wiley & Sons Ltd. (1995); Cid-Arregui (eds.), Viral Vectors: Essential Data, John Wiley & Sons Ltd. (1995); Cid-Arregui (eds.), Viral Vectors: Essential Data, John Wiley & Sons Ltd. (1996); Cid-Arregui (eds.), Viral Vectors: Essential Data, John Wiley & Sons Ltd. (1997); Cid-Arregui (eds.), Viral Vectors: Essential Data, John Wiley & Sons Ltd. (1998); Gold-Arregui (eds.), Viral Vectors: Essential Data, John Wiley & Sons Ltd. (1998); Gold-Arregui (eds.), Viral Vectors: Essential Data, John Wiley & Sons Ltd. (1998); Gold-Arregui (eds.), Viral Vectors: Essential Data, John Wiley & Sons Ltd. (1998); Gold-Arregui (eds.), Viral V

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Nucleic acid sequences may be expressed by operatively linking them to an expression control sequence in an appropriate expression vector and employing that expression vector to transform an appropriate unicellular host. Expression control sequences are sequences which control the transcription, post-transcriptional events and translation of nucleic acid sequences. Such operative linking of a nucleic sequence of this invention to an expression control sequence, of course, includes, if not already part of the nucleic acid sequence, the provision of a translation initiation codon, ATG or GTG, in the correct reading frame unstream of the nucleic acid sequence.

available commercially. Use of existing vectors and modifications thereof are well within

the skill in the art Thus, only basic features need be described here.

A wide variety of host/expression vector combinations may be employed in expressing the nucleic acid sequences of this invention. Useful expression vectors, for example, may consist of segments of chromosomal, non-chromosomal and synthetic nucleic acid sequences.

In one embodiment, prokaryotic cells may be used with an appropriate vector. Prokaryotic host cells are often used for cloning and expression. In a preferred embodiment, prokaryotic host cells include E. coli, Pseudomonas, Bacillus and Streptomyces. In a preferred embodiment, bacterial host cells are used to express the nucleic acid molecules of the instant invention. Useful expression vectors for bacterial hosts include bacterial plasmids, such as those from E. coli, Bacillus or Streptomyces,

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including pBluescript, pGEX-2T, pUC vectors, col E1, pCR1, pBR322, pMB9 and their derivatives, wider host range plasmids, such as RP4, phage DNAs, e.g., the numerous derivatives of phage lambda, e.g., NM989, \(\lambda\)GT10 and \(\lambda\)GT11, and other phages, e.g., \(\lambda\)H13 and filamentous single stranded phage DNA. Where \(\lambda\) c. is used as host, selectable markers are, analogously, chosen for selectivity in gram negative bacteria: e.g., twical markers confer resistance to antibiotics, such as ampicillin, terraeveline.

selectable markers are, analogously, chosen for selectivity in gram negative bacteria: e.g., typical markers confer resistance to antibiotics, such as ampicillin, tetracycline, chloramphenicol, kanamycin, streptomycin and zeocin; auxotrophic markers can also be used.

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In other embodiments, eukaryotic host cells, such as yeast, insect, mammalian or plant cells, may be used. Yeast cells, typically S. cerevisiae, are useful for eukaryotic genetic studies, due to the ease of targeting genetic changes by homologous recombination and the ability to easily complement genetic defects using recombinantly expressed proteins. Yeast cells are useful for identifying interacting protein components, e.g. through use of a two-hybrid system. In a preferred embodiment, yeast cells are useful for protein expression. Vectors of the present invention for use in yeast will typically, but not invariably, contain an origin of replication suitable for use in yeast and a selectable marker that is functional in yeast. Yeast vectors include Yeast Integrating plasmids (e.g., YIp5) and Yeast Replicating plasmids (the YRp and YEp series plasmids), Yeast Centromere plasmids (the YCp series plasmids), Yeast Artificial Chromosomes (YACs) which are based on yeast linear plasmids, denoted YLp, pGPD-2, 2µ plasmids and derivatives thereof, and improved shuttle vectors such as those described in Gietz et al., Gene, 74: 527-34 (1988) (YIplac, YEplac and YCplac). Selectable markers in yeast vectors include a variety of auxotrophic markers, the most common of which are (in Saccharomyces cerevisiae) URA3, HIS3, LEU2, TRP1 and LYS2, which complement specific auxotrophic mutations, such as ura3-52, his3-D1, leu2-D1, trp1-D1 and lys2-201.

Insect cells are often chosen for high efficiency protein expression. Where the host cells are from Spodoptera frugiperda, e.g., St9 and St21 cell lines, and expresSFTM cells (Protein Sciences Corp., Meriden, CT, USA), the vector replicative strategy is typically based upon the baculovirus life cycle. Typically, baculovirus transfer vectors are used to replace the wild-type AcMNPV polyhedrin gene with a heterologous gene of interest. Sequences that flank the polyhedrin gene in the wild-type genome are positioned 5' and 3' of the expression cassette on the transfer vectors. Following co-transfection with AcMNPV DNA, a homologous recombination event occurs between these sequences

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resulting in a recombinant virus carrying the gene of interest and the polyhedrin or p10 promoter. Selection can be based upon visual screening for lacZ fusion activity.

The host cells may also be mammalian cells, which are particularly useful for expression of proteins intended as pharmaceutical agents, and for screening of potential agonists and antagonists of a protein or a physiological pathway. Mammalian vectors intended for autonomous extrachromosomal replication will typically include a viral origin, such as the SV40 origin (for replication in cell lines expressing the large T-antigen, such as COS1 and COS7 cells), the papillomavirus origin, or the EBV origin for long term episomal replication (for use, e.g., in 293-EBNA cells, which constitutively express the EBV EBNA-1 gene product and adenovirus E1A). Vectors intended for integration, and thus replication as part of the mammalian chromosome, can, but need not, include an origin of replication functional in mammalian cells, such as the SV40 origin. Vectors based upon viruses, such as adenovirus, adeno-associated virus, vaccinia virus, and various mammalian retroviruses, will typically replicate according to the viral replicative strategy. Selectable markers for use in mammalian cells include, include but are not limited to, resistance to neomycin (G418), blasticidin, hygromycin and zeocin, and selection based upon the purine salvage pathway using HAT medium.

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Expression in mammalian cells can be achieved using a variety of plasmids, including pSV2, pBC12BI, and p91023, as well as lytic virus vectors (e.g., vaccinia virus, adeno virus, and baculovirus), episomal virus vectors (e.g., bovine papillomavirus), and retroviral vectors (e.g., murine retroviruses). Useful vectors for insect cells include baculoviral vectors and pVL 941.

Plant cells can also be used for expression, with the vector replicon typically derived from a plant virus (e.g., cauliflower mosaic virus, CaMV; tobacco mosaic virus, TMV) and selectable markers chosen for suitability in plants.

It is known that codon usage of different host cells may be different. For example, a plant cell and a human cell may exhibit a difference in codon preference for encoding a particular amino acid. As a result, human mRNA may not be efficiently translated in a plant, bacteria or insect host cell. Therefore, another embodiment of this invention is directed to codon optimization. The codons of the nucleic acid molecules of the invention may be modified to resemble, as much as possible, genes naturally contained within the host cell without altering the amino acid sequence encoded by the nucleic acid molecule.

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Any of a wide variety of expression control sequences may be used in these vectors to express the nucleic acid molecules of this invention. Such useful expression control sequences include the expression control sequences associated with structural genes of the foregoing expression vectors. Expression control sequences that control transcription include, e.g., promoters, enhancers and transcription termination sites. Expression control sequences in eukaryotic cells that control post-transcriptional events include splice donor and acceptor sites and sequences that modify the half-life of the transcribed RNA, e.g., sequences that direct poly(A) addition or binding sites for RNA-binding proteins. Expression control sequences that control translation include ribosome binding sites, sequences which direct targeted expression of the polypeptide to or within particular cellular compartments, and sequences in the 5' and 3' untranslated regions that modify the rate or efficiency of translation.

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Examples of useful expression control sequences for a prokaryote, e.g., E. coli, will include a promoter, often a phage promoter, such as phage lambda pL promoter, the tre promoter, a hybrid derived from the trp and lae promoters, the bacteriophage T7 promoter (in E. coli cells engineered to express the T7 polymerase), the TAC or TRC system, the major operator and promoter regions of phage lambda, the control regions of fd coat protein, and the araBAD operon. Prokaryotic expression vectors may further include transcription terminators, such as the aspA terminator, and elements that facilitate translation, such as a consensus ribosome binding site and translation termination codon, Schomer et al., Proc. Natl. Acad. Sci. USA 83: 8506-8510 (1986).

Expression control sequences for yeast cells, typically S. cerevisiae, will include a yeast promoter, such as the CYC1 promoter, the GAL1 promoter, the GAL10 promoter, ADH1 promoter, the promoters of the yeast α -mating system, or the GPD promoter, and will typically have elements that facilitate transcription termination, such as the transcription termination signals from the CYC1 or ADH1 gene.

Expression vectors useful for expressing proteins in mammalian cells will include a promoter active in mammalian cells. These promoters include, but are not limited to, those derived from mammalian viruses, such as the enhancer-promoter sequences from the immediate early gene of the human cytomegalovirus (CMV), the enhancer-promoter sequences from the Rous sarcoma virus long terminal repeat (RSV LTR), the enhancer-promoter from SV40 and the early and late promoters of adenovirus. Other expression control sequences include the promoter for 3-phosphoglycerate kinase or other glycolytic

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enzymes, the promoters of acid phosphatase. Other expression control sequences include those from the gene comprising the CSNA of interest. Often, expression is enhanced by incorporation of polyadenylation sites, such as the late SV40 polyadenylation site and the polyadenylation signal and transcription termination sequences from the bovine growth hormone (BGH) gene, and ribosome binding sites. Furthermore, vectors can include introns, such as intron II of rabbit β-globin gene and the SV40 splice elements.

Preferred nucleic acid vectors also include a selectable or amplifiable marker gene and means for amplifying the copy number of the gene of interest. Such marker genes are well-known in the art. Nucleic acid vectors may also comprise stabilizing sequences (e.g., ori- or ARS-like sequences and telomere-like sequences), or may alternatively be designed to favor directed or non-directed integration into the host cell genome. In a preferred embodiment, nucleic acid sequences of this invention are inserted in frame into an expression vector that allows high level expression of an RNA which encodes a protein comprising the encoded nucleic acid sequence of interest. Nucleic acid cloning and sequencing methods are well-known to those of skill in the art and are described in an assortment of laboratory manuals, including Sambrook (1989), supra, Sambrook (2000), supra; and Ausubel (1992), supra, Ausubel (1999), supra. Product information from manufacturers of biological, chemical and immunological reagents also provide useful information.

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Expression vectors may be either constitutive or inducible. Inducible vectors include either naturally inducible promoters, such as the tre promoter, which is regulated by the lac operon, and the pL promoter, which is regulated by tryptophan, the MMTV-LTR promoter, which is inducible by dexamethasone, or can contain synthetic promoters and/or additional elements that confer inducible control on adjacent promoters. Examples of inducible synthetic promoters are the hybrid Plac/ara-1 promoter and the PLtetO-1 promoter. The PltetO-1 promoter takes advantage of the high expression levels from the PL promoter of phage lambda, but replaces the lambda repressor sites with two copies of operator 2 of the Tn10 tetracycline resistance operon, causing this promoter to be tightly repressed by the Tet repressor protein and induced in response to tetracycline 30 (Tc) and Tc derivatives such as anhydrotetracycline. Vectors may also be inducible because they contain hormone response elements, such as the glucocorticoid response element (GRE) and the estrogen response element (ERE), which can confer hormone inducibility where vectors are used for expression in cells having the respective hormone

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receptors. To reduce background levels of expression, elements responsive to ecdysone, an insect hormone, can be used instead, with coexpression of the ecdysone receptor.

In one embodiment of the invention, expression vectors can be designed to fuse the expressed polypeptide to small protein tags that facilitate purification and/or visualization. Such tags include a polyhistidine tag that facilitates purification of the fusion protein by immobilized metal affinity chromatography, for example using NiNTA resin (Oiagen Inc., Valencia, CA, USA) or TALON™ resin (cobalt immobilized affinity chromatography medium, Clontech Labs, Palo Alto, CA, USA). The fusion protein can include a chitinbinding tag and self-excising intein, permitting chitin-based purification with self-removal of the fused tag (IMPACTTM system, New England Biolabs, Inc., Beverley, MA, USA). 10 Alternatively, the fusion protein can include a calmodulin-binding pentide tag, permitting purification by calmodulin affinity resin (Stratagene, La Jolla, CA, USA), or a specifically excisable fragment of the biotin carboxylase carrier protein, permitting purification of in vivo biotinylated protein using an avidin resin and subsequent tag removal (Promega, Madison, WI, USA). As another useful alternative, the polypeptides of the present 15 invention can be expressed as a fusion to glutathione-S-transferase, the affinity and specificity of binding to glutathione permitting purification using glutathione affinity resins, such as Glutathione-Superflow Resin (Clontech Laboratories, Palo Alto, CA. USA), with subsequent elution with free glutathione. Other tags include, for example, the Xpress epitope, detectable by anti-Xpress antibody (Invitrogen, Carlsbad, CA, USA), a 20 myc tag, detectable by anti-myc tag antibody, the V5 epitope, detectable by anti-V5 antibody (Invitrogen, Carlsbad, CA, USA), FLAG® epitope, detectable by anti-FLAG®

For secretion of expressed polypeptides, vectors can include appropriate sequences that encode secretion signals, such as leader peptides. For example, the pSecTag2 vectors (Invitrogen, Carlsbad, CA, USA) are 5.2 kb mammalian expression vectors that carry the secretion signal from the V-I2-C region of the mouse Ig kappa-chain for efficient secretion of recombinant proteins from a variety of mammalian cell lines.

antibody (Stratagene, La Jolla, CA, USA), and the HA epitope, detectable by anti-HA

antibody.

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Expression vectors can also be designed to fuse proteins encoded by the heterologous nucleic acid insert to polypeptides that are larger than purification and/or identification tags. Useful protein fusions include those that permit display of the encoded protein on the surface of a phage or cell, fusions to intrinsically fluorescent proteins, such 50

as those that have a green fluorescent protein (GFP)-like chromophore, fusions to the IgG Fc region, and fusions for use in two hybrid systems.

Vectors for phage display fuse the encoded polypeptide to, e.g., the gene III protein (pIII) or gene VIII protein (pVIII) for display on the surface of filamentous phage, such as M13. See Barbas et al., phage Display: A Laboratory Manual, Cold Spring Harbor Laboratory Press (2001); Kay et al. (eds.), <u>Phage Display of Peptides and Proteins: A Laboratory Manual</u>, Academic Press, Inc., (1996); Abelson et al. (eds.), <u>Combinatorial Chemistry</u> (Methods in Enzymology, Vol. 267) Academic Press (1996). Vectors for yeast display, e.g. the pYD1 yeast display vector (Invitrogen, Carlsbad, CA, USA), use the α-agglutinin yeast adhesion receptor to display recombinant protein on the surface of S. cerevisiae. Vectors for mammalian display, e.g., the pDisplayTM vector (Invitrogen, Carlsbad, CA, USA), target recombinant proteins using an N-terminal cell surface targeting signal and a C-terminal transmembrane anchoring domain of platelet derived growth factor receptor.

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A wide variety of vectors now exist that fuse proteins encoded by heterologous nucleic acids to the chromophore of the substrate-independent, intrinsically fluorescent green fluorescent protein from Aeguorea victoria ("GFP") and its variants. The GFP-like chromophore can be selected from GFP-like chromophores found in naturally occurring proteins, such as A. victoria GFP (GenBank accession number AAA27721), Renilla reniformis GFP, FP583 (GenBank accession no. AF168419) (DsRed), FP593 (AF272711). FP483 (AF168420), FP484 (AF168424), FP595 (AF246709), FP486 (AF168421), FP538 (AF168423), and FP506 (AF168422), and need include only so much of the native protein as is needed to retain the chromophore's intrinsic fluorescence. Methods for determining the minimal domain required for fluorescence are known in the art. See Li et al., J. Biol. Chem. 272; 28545-28549 (1997). Alternatively, the GFP-like chromophore can be selected from GFP-like chromophores modified from those found in nature. The methods for engineering such modified GFP-like chromophores and testing them for fluorescence activity, both alone and as part of protein fusions, are well-known in the art. See Heim et al., Curr. Biol. 6: 178-182 (1996) and Palm et al., Methods Enzymol. 302: 378-394 (1999). A variety of such modified chromophores are now commercially available and can readily be used in the fusion proteins of the present invention. These include EGFP ("enhanced GFP"), EBFP ("enhanced blue fluorescent protein"), BFP2, EYFP ("enhanced yellow

fluorescent protein"), ECFP ("enhanced cyan fluorescent protein") or Citrine. EGFP (see.

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e.g, Cormack et al., Gene 173: 33-38 (1996); United States Patent Nos. 6,090,919 and 5,804,387) is found on a variety of vectors, both plasmid and viral, which are available commercially (Clontech Labs, Palo Alto, CA, USA); EBFP is optimized for expression in mammalian cells whereas BFP2, which retains the original jellyfish codons, can be 5 expressed in bacteria (see, e.g., Heim et al., Curr. Biol. 6: 178-182 (1996) and Cormack et al., Gene 173: 33-38 (1996)). Vectors containing these blue-shifted variants are available from Clontech Labs (Palo Alto, CA, USA). Vectors containing EYFP, ECFP (see, e.g., Heim et al., Curr. Biol. 6: 178-182 (1996); Miyawaki et al., Nature 388: 882-887 (1997)) and Citrine (see, e.g., Heikal et al., Proc. Natl. Acad. Sci. USA 97: 11996-12001 (2000)) are also available from Clontech Labs. The GFP-like chromophore can also be drawn from other modified GFPs, including those described in United States Patent Nos. 6,124,128; 6,096,865; 6,090,919; 6,066,476; 6,054,321; 6,027,881; 5,968,750; 5,874,304; 5,804,387; 5,777,079; 5,741,668; and 5,625,048, the disclosures of which are incorporated herein by reference in their entireties. See also Conn (ed.), Green Fluorescent Protein (Methods in Enzymology, Vol. 302), Academic Press, Inc. (1999). The GFP-like chromophore of each of these GFP variants can usefully be included in the fusion proteins of the present invention.

Fusions to the IgG Fc region increase serum half life of protein pharmaceutical products through interaction with the FcRn receptor (also denominated the FcRp receptor and the Brambell receptor, FcRb), further described in International Patent Application nos. WO 97/43316. WO 97/34631. WO 96/32478. WO 96/18412.

For long-term, high-yield recombinant production of the polypeptides of the present invention, stable expression is preferred. Stable expression is readily achieved by integration into the host cell genome of vectors having selectable markers, followed by selection of these integrants. Vectors such as pUB6/V5-His A, B, and C (Invitrogen, Carlsbad, CA, USA) are designed for high-level stable expression of heterologous proteins in a wide range of mammalian tissue types and cell lines. pUB6/V5-His uses the promoter/enhancer sequence from the human ubiquitin C gene to drive expression of recombinant proteins: expression levels in 293, CHO, and NIH3T3 cells are comparable to levels from the CMV and human EF-1a promoters. The bsd gene permits rapid selection of stably transfected mammalian cells with the potent antibiotic blasticidin.

Replication incompetent retroviral vectors, typically derived from Moloney murine leukemia virus, also are useful for creating stable transfectants having integrated provirus.

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The highly efficient transduction machinery of retroviruses, coupled with the availability of a variety of packaging cell lines such as RetroPackTM PT 67, EcoPack^{ZTM}.293, AmphoPack-293, and GP2-293 cell lines (all available from Clontech Laboratories, Palo Alto, CA, USA) allow a wide host range to be infected with high efficiency; varying the multiplicity of infection readily adjusts the copy number of the integrated provirus.

Of course, not all vectors and expression control sequences will function equally well to express the nucleic acid molecules of this invention. Neither will all hosts function equally well with the same expression system. However, one of skill in the art may make a selection among these vectors, expression control sequences and hosts without undue experimentation and without departing from the scope of this invention. For example, in selecting a vector, the host must be considered because the vector must be replicated in it. The vector's copy number, the ability to control that copy number, the ability to control integration, if any, and the expression of any other proteins encoded by the vector, such as antibiotic or other selection markers, should also be considered. The present invention further includes host cells comprising the vectors of the present invention, either present episomally within the cell or integrated, in whole or in part, into the host cell chromosome. Among other considerations, some of which are described above, a host cell strain may be chosen for its ability to process the expressed polypeptide in the desired fashion. Such post-translational modifications of the polypeptide include, but are not limited to, acetylation, carboxylation, glycosylation, phosphorylation, lipidation, and acylation, and it is an aspect of the present invention to provide CSPs with such post-translational modifications.

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In selecting an expression control sequence, a variety of factors should also be considered. These include, for example, the relative strength of the sequence, its controllability, and its compatibility with the nucleic acid molecules of this invention, particularly with regard to potential secondary structures. Unicellular hosts should be selected by consideration of their compatibility with the chosen vector, the toxicity of the product coded for by the nucleic acid sequences of this invention, their secretion characteristics, their ability to fold the polypeptide correctly, their fermentation or culture requirements, and the ease of purification from them of the products coded for by the nucleic acid molecules of this invention.

The recombinant nucleic acid molecules and more particularly, the expression vectors of this invention may be used to express the polypeptides of this invention as

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recombinant polypeptides in a heterologous host cell. The polypeptides of this invention may be full-length or less than full-length polypeptide fragments recombinantly expressed from the nucleic acid molecules according to this invention. Such polypeptides include analogs, derivatives and muteins that may or may not have biological activity.

Vectors of the present invention will also often include elements that permit in vitro transcription of RNA from the inserted heterologous nucleic acid. Such vectors typically include a phage promoter, such as that from T7, T3, or SP6, flanking the nucleic acid insert. Often two different such promoters flank the inserted nucleic acid, permitting separate in vitro production of both sense and antisense strands.

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Transformation and other methods of introducing nucleic acids into a host cell (e.g., conjugation, protoplast transformation or fusion, transfection, electroporation, liposome delivery, membrane fusion techniques, high velocity DNA-coated pellets, viral infection and protoplast fusion) can be accomplished by a variety of methods which are well-known in the art (See, for instance, Ausubel, supra, and Sambrook et al., supra). Bacterial, yeast, plant or mammalian cells are transformed or transfected with an expression vector, such as a plasmid, a cosmid, or the like, wherein the expression vector comprises the nucleic acid of interest. Alternatively, the cells may be infected by a viral expression vector comprising the nucleic acid of interest. Depending upon the host cell, vector, and method of transformation used, transient or stable expression of the polypeptide will be constitutive or inducible. One having ordinary skill in the art will be express the protein constitutively or inducibly.

A wide variety of unicellular host cells are useful in expressing the DNA sequences of this invention. These hosts may include well-known eukaryotic and prokaryotic hosts, such as strains of, fungi, yeast, insect cells such as Spodoptera frugiperda (SF9), animal cells such as CHO, as well as plant cells in tissue culture. Representative examples of appropriate host cells include, but are not limited to, bacterial cells, such as E. coli, Caulobacter crescentus, Streptomyces species, and Salmonella typhimurium; yeast cells, such as Saccharomyces cerevisiae, Schizosaccharomyces pombe, Pichia pastoris, Pichia methanolica; insect cell lines, such as those from Spodoptera frugiperda — e.g., SF9 and SF21 cell lines, and expresSF™ cells (Protein Sciences Corp., Meriden, CT, USA) — Drosophila S2 cells, and Trichoplusia ni High Five® Cells (Invitrogen, Carlsbad, CA, USA); and mammalian cells. Typical mammalian cells include

BHK cells, BSC 1 cells, BSC 40 cells, BMT 10 cells, VERO cells, COS1 cells, COS7 cells, Chinese hamster ovary (CHO) cells, 3T3 cells, NIH 3T3 cells, 293 cells, HEPG2 cells, HeLa cells, L cells, MDCK cells, HEK293 cells, WI38 cells, murine ES cell lines (e.g., from strains 129/SV, C57/BL6, DBA-1, 129/SVJ), K562 cells, Jurkat cells, and BW5147 cells. Other mammalian cell lines are well-known and readily available from the American Type Culture Collection (ATCC) (Manassas, VA, USA) and the National Institute of General Medical Sciences (NIGMS) Human Genetic Cell Repository at the Coriell Cell Repositories (Camden, NJ, USA). Cells or cell lines derived from colon are particularly preferred because they may provide a more native post-translational processing. Particularly preferred are human colon cells.

Particular details of the transfection, expression and purification of recombinant proteins are well documented and are understood by those of skill in the art. Further details on the various technical aspects of each of the steps used in recombinant production of foreign genes in bacterial cell expression systems can be found in a number of texts and laboratory manuals in the art. See, e.g., Ausubel (1992), supra, Ausubel (1999), supra, Sambrook (1989), supra, and Sambrook (2001), supra.

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Methods for introducing the vectors and nucleic acid molecules of the present invention into the host cells are well-known in the art; the choice of technique will depend primarily upon the specific vector to be introduced and the host cell chosen.

Nucleic acid molecules and vectors may be introduced into prokaryotes, such as *E. coli*, in a number of ways. For instance, phage lambda vectors will typically be packaged using a packaging extract (e.g., Gigapack® packaging extract, Stratagene, La Jolla, CA, USA), and the packaged virus used to infect *E. coli*.

Plasmid vectors will typically be introduced into chemically competent or electrocompetent bacterial cells. *E. coli* cells can be rendered chemically competent by treatment, *e.g.*, with CaCl₂, or a solution of Mg²⁺, Mn²⁺, Ca²⁺, Rb⁺ or K⁺, dimethyl sulfoxide, dithiothreitol, and hexamine cobalt (III), Hanahan, *J. Mol. Biol.* 166(4):557-80 (1983), and vectors introduced by heat shock. A wide variety of chemically competent strains are also available commercially (*e.g.*, Epicurian Coli® XL10-Gold®

0 Ultracompetent Cells (Stratagene, La Jolla, CA, USA); DH5α competent cells (Clontech Laboratories, Palo Alto, CA, USA); and TOP10 Chemically Competent E. coli Kit (Invitrogen, Carlsbad, CA, USA)). Bacterial cells can be rendered electrocompetent to take up exogenous DNA by electroporation by various pre-pulse treatments; vectors are introduced by electroporation followed by subsequent outgrowth in selected media. An extensive series of protocols is provided by BioRad (Richmond, CA, USA).

Vectors can be introduced into yeast cells by spheroplasting, treatment with lithium salts, electroporation, or protoplast fusion. Spheroplasts are prepared by the action of hydrolytic enzymes such as a snail-gut extract, usually denoted Glusulase or Zymolyase, or an enzyme from Arthrobacter Inteus to remove portions of the cell wall in the presence of osmotic stabilizers, typically 1 M sorbitol. DNA is added to the spheroplasts, and the mixture is co-precipitated with a solution of polyethylene glycol (PEG) and Ca²⁺. Subsequently, the cells are resuspended in a solution of sorbitol, mixed with molten agar and then layered on the surface of a selective plate containing sorbitol.

For lithium-mediated transformation, yeast cells are treated with lithium acetate to permeabilize the cell wall, DNA is added and the cells are co-precipitated with PEG. The cells are exposed to a brief heat shock, washed free of PEG and lithium acetate, and subsequently spread on plates containing ordinary selective medium. Increased frequencies of transformation are obtained by using specially-prepared single-stranded carrier DNA and certain organic solvents. Schiestl et al., Curr. Genet. 16(5-6): 339-46 (1989).

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For electroporation, freshly-grown yeast cultures are typically washed, suspended in an osmotic protectant, such as sorbitol, mixed with DNA, and the cell suspension pulsed in an electroporation device. Subsequently, the cells are spread on the surface of plates containing selective media. Becker et al., Methods Enzymol. 194: 182-187 (1991). The efficiency of transformation by electroporation can be increased over 100-fold by using PEG, single-stranded carrier DNA and cells that are in late log-phase of growth. Larger constructs, such as YACs, can be introduced by protoplast fusion.

Mammalian and insect cells can be directly infected by packaged viral vectors, or transfected by chemical or electrical means. For chemical transfection, DNA can be coprecipitated with CaPO₄ or introduced using liposomal and nonliposomal lipid-based agents. Commercial kits are available for CaPO₄ transfection (CalPhos™ Mammalian Transfection Kit, Clontech Laboratories, Palo Alto, CA, USA), and lipid-mediated transfection can be practiced using commercial reagents, such as LIPOFECTAMINE™ 2000, LIPOFECTAMINE™ Reagent, CELLFECTIN® Reagent, and LIPOFECTIN® Reagent (Invitrogen, Carlsbad, CA, USA), DOTAP Liposomal Transfection Reagent, FuGENE 6, X-tremeGENE 02. DOSPER. (Roche Molecular Biochemicals, Indiananolis,

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IN USA), Effectene™, PolyFect®, Superfect® (Qiagen, Inc., Valencia, CA, USA).

Protocols for electroporating mammalian cells can be found in, for example,; Norton et al. (eds.), Gene Transfer Methods: Introducing DNA into Living Cells and Organisms,

BioTechniques Books, Eaton Publishing Co. (2000). Other transfection techniques include transfection by particle bombardment and microinjection. See, e.g., Cheng et al., Proc. Natl. Acad. Sci. USA 90(10): 4455-9 (1993); Yang et al., Proc. Natl. Acad. Sci. USA 87(24): 9568-72 (1990).

Production of the recombinantly produced proteins of the present invention can optionally be followed by purification.

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Purification of recombinantly expressed proteins is now well within the skill in the art and thus need not be detailed here. See, e.g., Thomer et al. (eds.), <u>Applications of Chimeric Genes and Hybrid Proteins. Part A: Gene Expression and Protein Purification</u> (Methods in Enzymology, Vol. 326), Academic Press (2000); Harbin (ed.), <u>Cloning. Gene Expression and Protein Purification</u>: <u>Experimental Procedures and Process Rationale</u>.

Oxford Univ. Press (2001); Marshak et al., Strategies for Protein Purification and Characterization: A Laboratory Course Manual, Cold Spring Harbor Laboratory Press (1996); and Roe (ed.), Protein Purification Applications. Oxford University Press (2001).

Briefly, however, if purification tags have been fused through use of an expression vector that appends such tag, purification can be effected, at least in part, by means appropriate to the tag, such as use of immobilized metal affinity chromatography for polyhistidine tags. Other techniques common in the art include ammonium sulfate fractionation, immunoprecipitation, fast protein liquid chromatography (FPLC), high performance liquid chromatography (HPLC), and preparative gel electrophoresis.

Polypeptides, including Fragments Muteins, Homologous Proteins, Allelic Variants,
Analogs and Derivatives

Another aspect of the invention relates to polypeptides encoded by the nucleic acid molecules described herein. In a preferred embodiment, the polypeptide is a colon specific polypeptide (CSP). In an even more preferred embodiment, the polypeptide comprises an amino acid sequence of SEQ ID NO:59-82 or is derived from a polypeptide having the amino acid sequence of SEQ ID NO: 101-190. A polypeptide as defined herein may be produced recombinantly, as discussed supra, may be isolated from a cell that naturally expresses the protein, or may be chemically synthesized following the teachings

of the specification and using methods well-known to those having ordinary skill in the art.

Polypeptides of the present invention may also comprise a part or fragment of a CSP. In a preferred embodiment, the fragment is derived from a polypeptide having an amino acid sequence selected from the group consisting of SEQ ID NO: 101-190. Polypeptides of the present invention comprising a part or fragment of an entire CSP may or may not be CSPs. For example, a full-length polypeptide may be colon-specific, while a fragment thereof may be found in other tissues as well as in colon. A polypeptide that is not a CSP, whether it is a fragment, analog, mutein, homologous protein or derivative, is nevertheless useful, especially for immunizing animals to prepare anti-CSP antibodies. In a preferred embodiment, the part or fragment is a CSP. Methods of determining whether a polypeptide of the present invention is a CSP are described infrac.

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Polypeptides of the present invention comprising fragments of at least 6 contiguous amino acids are also useful in mapping B cell and T cell epitopes of the reference protein. See, e.g., Geysen et al., Proc. Natl. Acad. Sci. USA 81: 3998-4002 (1984) and United States Patent Nos. 4,708,871 and 5,595,915, the disclosures of which are incorporated herein by reference in their entireties. Because the fragment need not itself be immunogenic, part of an immunodominant epitope, nor even recognized by native antibody, to be useful in such epitope mapping, all fragments of at least 6 amino acids of a polypeptide of the present invention have utility in such a study.

Polypeptides of the present invention comprising fragments of at least 8 contiguous amino acids, often at least 15 contiguous amino acids, are useful as immunogens for raising antibodies that recognize polypeptides of the present invention. See, e.g., Lerner, Nature 299: 592-596 (1982); Shinnick et al., Annu. Rev. Microbiol. 37: 25 425-46 (1983); Sutcliffe et al., Science 219: 660-6 (1983). As further described in the above-cited references, virtually all 8-mers, conjugated to a carrier, such as a protein, prove immunogenic and are capable of eliciting antibody for the conjugated peptide; accordingly, all fragments of at least 8 amino acids of the polypeptides of the present invention have utility as immunogens.

Polypeptides comprising fragments of at least 8, 9, 10 or 12 contiguous amino acids are also useful as competitive inhibitors of binding of the entire polypeptide, or a portion thereof, to antibodies (as in epitope mapping), and to natural binding partners, such as subunits in a multimeric complex or to receptors or ligands of the subject protein; this competitive inhibition permits identification and separation of molecules that bind specifically to the polypeptide of interest. See United States Patent Nos. 5,539,084 and 5,783,674, incorporated herein by reference in their entireties.

The polypeptide of the present invention thus preferably is at least 6 amino acids in length, typically at least 8, 9, 10 or 12 amino acids in length, and often at least 15 amino acids in length. Often, the polypeptide of the present invention is at least 20 amino acids in length, even 25 amino acids, 30 amino acids, 35 amino acids, or 50 amino acids or more in length. Of course, larger polypeptides having at least 75 amino acids, 100 amino acids, or even 150 amino acids are also useful, and at times preferred.

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One having ordinary skill in the art can produce fragments by truncating the nucleic acid molecule, e.g., a CSNA, encoding the polypeptide and then expressing it recombinantly. Alternatively, one can produce a fragment by chemically synthesizing a portion of the full-length polypeptide. One may also produce a fragment by enzymatically cleaving either a recombinant polypeptide or an isolated naturally-occurring polypeptide. Methods of producing polypeptide fragments are well-known in the art. See, e.g., Sambrook (1989), supra; Sambrook (2001), supra; Ausubel (1992), supra; and Ausubel (1999), supra. In one embodiment, a polypeptide comprising only a fragment, preferably a fragment of a CSP, may be produced by chemical or enzymatic cleavage of a CSP polypeptide. In a preferred embodiment, a polypeptide fragment is produced by expressing a nucleic acid molecule of the present invention encoding a fragment, preferably of a CSP, in a host cell.

Polypeptides of the present invention are also inclusive of mutants, fusion proteins, homologous proteins and allelic variants.

A mutant protein, or mutein, may have the same or different properties compared
to a naturally-occurring polypeptide and comprises at least one amino acid insertion,
duplication, deletion, rearrangement or substitution compared to the amino acid sequence
of a native polypeptide. Small deletions and insertions can often be found that do not alter
the function of a protein. Muteins may or may not be colon-specific. Preferably, the
mutein is colon-specific. More preferably the mutein is a polypeptide that comprises at
least one amino acid insertion, duplication, deletion, rearrangement or substitution
compared to the amino acid sequence of SEQ ID NO: 101-190. Accordingly, in a
preferred embodiment, the mutein is one that exhibits at least 50% sequence identity, more
preferably at least 60% sequence identity, even more preferably at least 70%, yet more

preferably at least 80% sequence identity to a CSP comprising an amino acid sequence of SEO ID NO: 101-190. In a yet more preferred embodiment, the mutein exhibits at least 85%, more preferably 90%, even more preferably 95% or 96%, and yet more preferably at least 97%, 98%, 99% or 99.5% sequence identity to a CSP comprising an amino acid sequence of SEQ ID NO: 101-190.

A mutein may be produced by isolation from a naturally-occurring mutant cell, tissue or organism. A mutein may be produced by isolation from a cell, tissue or organism that has been experimentally mutagenized. Alternatively, a mutein may be produced by chemical manipulation of a polypeptide, such as by altering the amino acid residue to another amino acid residue using synthetic or semi-synthetic chemical techniques. In a preferred embodiment, a mutein is produced from a host cell comprising a mutated nucleic acid molecule compared to the naturally-occurring nucleic acid molecule. For instance, one may produce a mutein of a polypeptide by introducing one or more mutations into a nucleic acid molecule of the invention and then expressing it recombinantly. These mutations may be targeted, in which particular encoded amino acids are altered, or may be untargeted, in which random encoded amino acids within the polypeptide are altered. Muteins with random amino acid alterations can be screened for a particular biological activity or property, particularly whether the polypeptide is colon-specific, as described below. Multiple random mutations can be introduced into the gene by methods wellknown to the art, e.g., by error-prone PCR, shuffling, oligonucleotide-directed mutagenesis, assembly PCR, sexual PCR mutagenesis, in vivo mutagenesis, cassette mutagenesis, recursive ensemble mutagenesis, exponential ensemble mutagenesis and sitespecific mutagenesis. Methods of producing muteins with targeted or random amino acid alterations are well-known in the art. See, e.g., Sambrook (1989), supra; Sambrook 25 (2001), supra; Ausubel (1992), supra; and Ausubel (1999), as well as United States Patent No. 5,223,408, which is herein incorporated by reference in its entirety.

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The invention also contemplates polypeptides that are homologous to a polypeptide of the invention. In a preferred embodiment, the polypeptide is homologous to a CSP. In an even more preferred embodiment, the polypeptide is homologous to a CSP selected from the group having an amino acid sequence of SEO ID NO: 101-190. By homologous polypeptide it is meant s one that exhibits significant sequence identity to a CSP, preferably a CSP having an amino acid sequence of SEQ ID NO: 101-190. By significant sequence identity it is meant that the homologous polypeptide exhibits at least

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50% sequence identity, more preferably at least 60% sequence identity, even more preferably at least 70%, yet more preferably at least 80% sequence identity to a CSP comprising an amino acid sequence of SEQ ID NO: 101-190. More preferred are homologous polypeptides exhibiting at least 85%, more preferably 90%, even more preferably 95% or 96%, and yet more preferably at least 97% or 98% sequence identity to a CSP comprising an amino acid sequence of SEQ ID NO: 101-190. Most preferably, the homologous polypeptide exhibits at least 99%, more preferably 99.5%, even more preferably 99.6%, 99.7%, 99.8% or 99.9% sequence identity to a CSP comprising an amino acid sequence of SEQ ID NO: 101-190. In a preferred embodiment, the amino acid substitutions of the homologous polypeptide are conservative amino acid substitutions as discussed above.

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Homologous polypeptides of the present invention also comprise polypeptide encoded by a nucleic acid molecule that selectively hybridizes to a CSNA or an antisense sequence thereof. In this embodiment, it is preferred that the homologous polypeptide be encoded by a nucleic acid molecule that hybridizes to a CSNA under low stringency, moderate stringency or high stringency conditions, as defined herein. More preferred is a homologous polypeptide encoded by a nucleic acid sequence which hybridizes to a CSNA selected from the group consisting of SEQ ID NO: 1-100 or a homologous polypeptide encoded by a nucleic acid molecule that hybridizes to a nucleic acid molecule that encodes a CSP, preferably an CSP of SEQ ID NO:59-82 under low stringency, moderate stringency or high stringency conditions, as defined herein.

Homologous polypeptides of the present invention may be naturally-occurring and derived from another species, especially one derived from another primate, such as chimpanzee, gorilla, rhesus macaque, or baboon, wherein the homologous polypeptide comprises an amino acid sequence that exhibits significant sequence identity to that of SEQ ID NO: 101-190. The homologous polypeptide may also be a naturally-occurring polypeptide from a human, when the CSP is a member of a family of polypeptides. The homologous polypeptide may also be a naturally-occurring polypeptide derived from a non-primate, mammalian species, including without limitation, domesticated species, e.g., dog, cat, mouse, rat, rabbit, guinea pig, hamster, cow, horse, goat or pig. The homologous polypeptide may also be a naturally-occurring polypeptide derived from a non-mammalian species, such as birds or reptiles. The naturally-occurring homologous protein may be isolated directly from humans or other species. Alternatively, the nucleic acid molecule

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encoding the naturally-occurring homologous polypeptide may be isolated and used to express the homologous polypeptide recombinantly. The homologous polypeptide may also be one that is experimentally produced by random mutation of a nucleic acid molecule and subsequent expression of the nucleic acid molecule. Alternatively, the homologous polypeptide may be one that is experimentally produced by directed mutation of one or more codons to alter the encoded amino acid of a CSP. In a preferred embodiment, the homologous polypeptide encodes a polypeptide that is a CSP.

Relatedness of proteins can also be characterized using a second functional test, the ability of a first protein competitively to inhibit the binding of a second protein to an antibody. It is, therefore, another aspect of the present invention to provide isolated polyeptide not only identical in sequence to those described with particularity herein, but also to provide isolated polyeptide ("cross-reactive proteins") that competitively inhibit the binding of antibodies to all or to a portion of various of the isolated polypeptides of the present invention. Such competitive inhibition can readily be determined using immunoassays well-known in the art.

As discussed above, single nucleotide polymorphisms (SNPs) occur frequently in eukaryotic genomes, and the sequence determined from one individual of a species may differ from other allelic forms present within the population. Thus, polypeptides of the present invention are also inclusive of those encoded by an allelic variant of a nucleic acid molecule encoding a CSP. In this embodiment, it is preferred that the polypeptide be encoded by an allelic variant of a gene that encodes a polypeptide having the amino acid sequence selected from the group consisting of SEQ ID NO: 101-190. More preferred is that the polypeptide be encoded by an allelic variant of a gene that has the nucleic acid sequence selected from the group consisting of SEO ID NO: 1-100.

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Polypeptides of the present invention are also inclusive of derivative polypeptides encoded by a nucleic acid molecule according to the instant invention. In this embodiment, it is preferred that the polypeptide be a CSP. Also preferred are derivative polypeptides having an amino acid sequence selected from the group consisting of SEQ ID NO: 101-190 and which has been acetylated, carboxylated, phosphorylated, glycosylated or ubiquitinated. In another preferred embodiment, the derivative has been labeled with, e.g., radioactive isotopes such as ¹²⁵I, ²³P, ³⁵S, and ³H. In another preferred

embodiment, the derivative has been labeled with fluorophores, chemiluminescent agents.

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enzymes, and antiligands that can serve as specific binding pair members for a labeled ligand.

Polypeptide modifications are well-known to those of skill and have been described in great detail in the scientific literature. Several particularly common modifications, glycosylation, lipid attachment, sulfation, gamma-carboxylation of glutamic acid residues, hydroxylation and ADP-ribosylation, for instance, are described in most basic texts, such as, for instance Creighton, <u>Protein Structure and Molecular Properties</u>, 2nd ed., W. H. Freeman and Company (1993). Many detailed reviews are available on this subject, such as, for example, those provided by Wold, in Johnson (ed.), <u>Posttranslational Covalent Modification of Proteins</u>, pgs. 1-12, Academic Press (1983); Seiffer et al., Meth. Enzymol. 182: 626-646 (1990) and Rattan et al., Ann. N.Y. Acad. Sci. 663: 48-62 (1992).

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One may determine whether a polypeptide of the invention is likely to be post-translationally modified by analyzing the sequence of the polypeptide to determine if there are peptide motifs indicative of sites for post-translational modification. There are a number of computer programs that permit prediction of post-translational modifications. See, e.g., www.expasy.org (accessed August 31, 2001), which includes PSORT, for prediction of protein sorting signals and localization sites, SignalP, for prediction of signal peptide cleavage sites, MITOPROT and Predotar, for prediction of mitochondrial targeting sequences, NetOGlyc, for prediction of type O-glycosylation sites in mammalian proteins, big-PI Predictor and DGPI, for prediction of prenylation-anchor and cleavage sites, and NetPhos, for prediction of Ser, Thr and Tyr phosphorylation sites in eukaryotic proteins. Other computer programs, such as those included in GCG, also may be used to determine post-translational modification peptide motifs.

General examples of types of post-translational modifications include, but are not limited to: (Z)-dehydrobutyrine; 1-chondroitin sulfate-L-aspartic acid ester; 1¹-glycosyl-L-tryptophan; 1¹-phospho-L-histidine; 1-thioglycine; 2¹-(S-L-cysteinyl)-L-histidine; 2¹-(3-L-cysteinyl)-L-histidine; 2¹-(3-L-cysteinyl)-L-tryptophan; 2-methyl-L-glutamine; 2-oxobutanoic acid; 2-pyrrolidone carboxylic acid; 3¹-(1¹-L-histidyl)-L-tyrosine; 3¹-(8alpha-FAD)-L-histidine; 3¹-(S-L-cysteinyl)-L-tyrosine; 3¹, 3″,5¹-triiodo-L-thyronine; 3¹-4¹-phospho-L-tyrosine; 3-hydroxy-L-proline; 3'-methyl-L-histidine; 3-methyl-L-hatthionine; 3¹-phospho-L-histidine; 4¹-(L-tryptophan)-L-tryptophyl quinone; 42 N-cysteinyl-glycosylphosphatidylinositolethanolamine; 43 -(T-L-histidyl)-L-tyrosine; 4-

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hydroxy-L-arginine; 4-hydroxy-L-lysine; 4-hydroxy-L-proline; 5'-{N6-L-lysine}-Ltopaquinone; 5-hydroxy-L-lysine; 5-methyl-L-arginine; alpha-l-microglobulin-Ig alpha complex chromophore; bis-L-cysteinyl bis-L-histidino dfiron disulfide; bis-L-cysteinyl-L-N3'-histidino-L-serinyl tetrairon' tetrasulfide; chondroitin sulfate D-glucuronyl-D-

- 5 galactosyl-D-galactosyl-D-xylosyl-L-serine; D-alanine; D-allo-isoleucine; D-asparagine; dehydroalanine; dehydrotyrosine; dermatan 4-sulfate D-glueuronyl-D-galactosyl-D-xylosyl-L-serine; D-glucuronyl-N-glycine; dipyrrolylmethanemethyl-L-cysteine; D-leucine; D-methionine; D-phenylalanine; D-serine; D-tryptophan; glycine amide; glycine oxazolecarboxylic acid; glycine thiazolecarboxylic acid; heme P450-bis-L-cysteine; L-tyrosine; heme-bis-L-cysteine; hemediol-L-aspartyl ester-L-glutamyl ester;
 - cysteme-L-tyrosine, heme-0is-L-cysteme, hemedioi-L-aspartyl ester-L-glutamyl ester, hemedioi-L-aspartyl ester-L-glutamyl ester-L-methionine sulfonium; heme-L-cysteine; heme-L-histidine; heparan sulfate D-glucuronyl-D-galactosyl-D-zylosyl-L-serine; heme P450-bis-L-cysteine-L-lysine; hexakis-L-cysteinyl hexairon hexasulfide; keratan sulfate D-glucuronyl-D-galactosyl-D-zylosyl-L-threonine; L
- 15 oxoalanine- lactic acid; L phenyllactic acid; l'-(8alpha-FAD)-L-histidine; L-2'.4',5'-topaquinone; L-3',4'-dihydroxyphenylalanine; L-3'.4'.5'-trihydroxyphenylalanine; L-4'-bromophenylalanine; L-6'-bromotyptophan; L-alanine amide; L-alanyl imidazolinone glycine; L-allysine; L-arginine amide; L-asparaigine amide; L-aspartic 4-phosphoric anhydride; L-aspartic acid 1-amide; L-beta-methylthioaspartic acid; L-bromohistidine; L-cirulline; L-cvsteine amide; L-evsteine glutathione disulfide: L-cvsteine methyl disulfide:
- L-cysteine methyl ester; L-cysteine oxazolecarboxylic acid; L-cysteine methyl ester; L-cysteine persulfide; L-cysteine sulfenic acid; L-cysteine sulfinic acid; L-cysteine persulfide; L-cysteine sulfinic acid; L-cysteine sulfinic acid; L-cysteinel miazolecarboxylic acid; L-cysteinyl homocitryl molybdonum-heptairon-nonasulfide; L-cysteinyl midazolinone glycine; L-cysteinyl molybdonterin guanine dinucleotide: L-cystein; L-cysteinyl molybdonterin guanine dinucleotide: L-cystein; L-cysteinyl molybdonterin guanine dinucleotide: L-cysteine L-cysteine sulfenic acid; L-cysteine sulfenic
- hydroxyasparagine; L-erythro-beta-hydroxyaspartic acid; L-gamma-carboxyglutamic acid; L-glutamic acid 1-amide; L-glutamyl cacid 5-methyl ester; L-glutamine amide; L-glutamyl 5-glycerylphosphorylethanolarmine; L-histidine amide; L-isoglutamyl-polyglycine; L-isoglut
- 30 amide; L-lysine thiazolecarboxylic acid; L-lysinoalanine; L-methionine amide; L-methionine sulfone; L-phenyalanine thiazolecarboxylic acid; L-phenylalanine amide; L-proline amide; L-selenocysteine; L-selenocysteinyl molybdopterin guanine dinucleotide; L-serine amide; L-serine thiazolecarboxylic acid; L-seryl imidazolinone glycine; L-T-

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bromophenylalanine; L-T-bromophenylalanine; L-threonine amide; L-thyroxine; Ltryptophan amide; L-tryptophyl quinone; L-tyrosine amide; L-valine amide; mesolanthionine; N-(L-glutamyl)-L-tyrosine; N-(L-isoaspartyl)-glycine; N-(L-isoaspartyl)-Lcysteine; N,N,N-trimethyl-L-alanine; N,N-dimethyl-L-proline; N2-acetyl-L-lysine; N2succinvl-L-tryptophan; N4-(ADP-ribosyl)-L-asparagine; N4-glycosyl-L-asparagine; N4hydroxymethyl-L-asparagine: N4-methyl-L-asparagine: N5-methyl-L-glutamine: N6- 1 carboxyethyl-L-lysine; N6-(4-amino hydroxybutyl)-L-lysine; N6-(L-isoglutamyl)-Llysine: N6-(phospho-5'-adenosine)-L-lysine: N6-(phospho-5'-guanosine)-L-tysine: N6,N6,N6-trimethyl-L-lysine; N6,N6-dimethyl-L-lysine; N6-acetyl-L-lysine; N6-biotinyl-L-Ivsine; N6-carboxy-L-Ivsine; N6-formyl-L-Ivsine; N6-glycyl-L-Ivsine; N6-lipoyl-Llysine: N6-methyl-L-lysine: N6-methyl-N6-poly(N-methyl-propylamine)-L-lysine: N6mureinyl-L-lysine; N6-myristoyl-L-lysine; N6-palmitoyl-L-lysine; N6-pyridoxal phosphate-L-lysine; N6-pyruvic acid 2-iminyl-L-lysine; N6-retinal-L-lysine; Nacetylglycine; N-acetyl-L.glutariaine; N-acetyl-L-alanine; N-acetyl-L-aspartic acid; Nacetyl-L-cysteine; N-acetyl-L-glutamic acid; N-acetyl-L-isoleucine; N-acetyl-Lmethionine; N-acetyl-L-proline; N-acetyl-L-serine; N-acetyl-L-threonine; N-acetyl-Ltyrosine; N-acetyl-L-valine; N-alanyl-glycosylphosphatidylinositolethanolamine; Nasparaginyl-glycosylphosphatidylinositolethanolarnine; N-aspartylglycosylphosphatidylinositolethanolanline; N-formylglycine; N-formyl-L-methionine; Nglycyl-glycosylphosphatidylinositolethanolarnine; N-L-glutamyl-poly-L-glutamic acid; Nmethylglycine; N-methyl-L-alanine; N-methyl-L-methionine; N-methyl-L-phenylalanine; N-myristoyl-glycine; N-palmitoyl-L-cysteine; N-pyruvic acid 2-iminyl-L-cysteine; Npyruvic acid 2-iminyl-L-valine: N-seryl-glycosylphosphatidylinositolethanolarnine: Nseryl-glycosycsphingolipidinositolethanolamine; O-(ADP-ribosyl)-L-serine; O-(phospho-5'-adenosine)-L-threonine; O-(phospho-5'-DNA)-L-serine; O-(phospho-5'-DNA)-Lthreonine; O-(phospho-5'rRNA)-L-serine; O-(phosphoribosyl dephospho-coenzyme A)-Lserine: O-(sn-l-glycerophosphoryl)-L-serine: O4'-(8alpha-FAD)-L-tyrosine: O4'-(phospho-5'-adenosine)-L-tyrosine; O4'-(phospho-5'-DNA)-L-tyrosine; O4'-(phospho-5'-RNA)-Ltyrosine; O4'-(phospho-5'-uridine)-L-tyrosine; O4-glycosyl-L-hydroxyproline; O4'-

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30 glycosyl-L-tyrosine; O4'-sulfo-L-tyrosine; O5-glycosyl-L-hydroxylysine; O-glycosyl-Lserine; O-glycosyl-L-threonine; omega-N-(ADP-ribosyl)-L-arginine; omega-N,omega-N'dimethyl-L-arginine; omega-N-methyl-L-arginine; omega-Nomega-N-dimethyl-Larginine; omega-N-phospho-L-arginine; O'octanoyl-L-serine.; O-palmitoyl-L-serine; O-

palmitovl-L-threonine; O-phospho-L-serine; O-phospho-L-threonine; Ophosphopantetheine-L-serine; phycoerythrobilin-bis-L-cysteine; phycourobilin-bis-Lcysteine; pyrroloquinoline quinone; pyruvic acid; S hydroxycinnamyl-L-cysteine; S-(2aminovinyl) methyl-D-eysteine; S-(2-aminovinyl)-D-cysteine; S-(6-FW-L-cysteine; S-(8alpha-FAD)-L-cysteine; S-(ADP-ribosyl)-L-cysteine; S-(L-isoglutamyl)-L-cysteine; S-12-hydroxyfarnesyl-L-cysteine; S-acetyl-L-cysteine; S-diacylglycerol-L-cysteine; Sdiphytanylglycerot diether-L-cysteine; S-farnesyl-L-cysteine; S-geranylgeranyl-Lcysteine: S-glycosyl-L-cysteine: S-glycyl-L-cysteine: S-methyl-L-cysteine: S-nitrosyl-Lcysteine; S-palmitoyl-L-cysteine; S-phospho-L-cysteine; S-phycobiliviolin-L-cysteine; S-10 phycocyanobilin-L-cysteine; S-phycoerythrobilin-L-cysteine; S-phytochromobilin-Lcysteine: S-selenyl-L-cysteine: S-sulfo-L-cysteine: tetrakis-L-cysteinyl diiron disulfide: tetrakis-L-cysteinyl iron; tetrakis-L-cysteinyl tetrairon tetrasulfide; trans-2,3-cis 4dihydroxy-L-proline; tris-L-cysteinyl triiron tetrasulfide; tris-L-cysteinyl triiron trisulfide; tris-L-cysteinyl-L-aspartato tetrairon tetrasulfide: tris-L-cysteinyl-L-cysteine persulfido-15 bis-L-glutamato-L-histidino tetrairon disulfide trioxide; tris-L-cysteinyl-L-N3'-histidino tetrairon tetrasulfide; tris-L-cysteinyl-L-Nl'-histidino tetrairon tetrasulfide; and tris-Lcysteinyl-L-serinyl tetrairon tetrasulfide.

Additional examples of PTMs may be found in web sites such as the Delta Mass database based on Krishna, R. G. and F. Wold (1998). Posttranslational Modifications. Proteins - Analysis and Design. R. H. Angeletti. San Diego, Academic Press. 1: 121-206.; Methods in Enzymology, 193, J.A. McClosky (ed) (1990), pages 647-660; Methods in Protein Sequence Analysis edited by Kazutomo Imahori and Fumio Sakiyama, Plenum Press. (1993) "Post-translational modifications of proteins" R.G. Krishna and F. Wold pages 167-172; "GlycoSuiteDB: a new curated relational database of glycoprotein glycan 25 structures and their biological sources" Cooper et al. Nucleic Acids Res. 29: 332-335 (2001) "O-GLYCBASE version 4.0: a revised database of O-glycosylated proteins" Gupta et al. Nucleic Acids Research, 27: 370-372 (1999); and "PhosphoBase, a database of phosphorvlation sites: release 2.0.", Kreegipuu et al.Nucleic Acids Res 27(1):237-239 (1999) see also, WO 02/21139A2.

Tumorigenesis is often accompanied by alterations in the post-translational modifications of proteins. Thus, in another embodiment, the invention provides polypeptides from cancerous cells or tissues that have altered post-translational modifications compared to the post-translational modifications of polypeptides from

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normal cells or tissues. A number of altered post-translational modifications are known. One common alteration is a change in phosphorylation state, wherein the polypeptide from the cancerous cell or tissue is hyperphosphorylated or hypophosphorylated compared to the polypeptide from a normal tissue, or wherein the polypeptide is phosphorylated on different residues than the polypeptide from a normal cell. Another common alteration is a change in glycosylation state, wherein the polypeptide from the cancerous cell or tissue has more or less glycosylation than the polypeptide from a normal tissue, and/or wherein the polypeptide from the cancerous cell or tissue has a different type of glycosylation than the polypeptide from a noncancerous cell or tissue has a different type of glycosylation than the polypeptide from a noncancerous cell or tissue has a different type of glycosylation may be critical because carbohydrate-protein and carbohydrate-carbohydrate interactions are important in cancer cell progression, dissemination and invasion. See, e.g., Barchi, Curr. Pharm. Des. 6: 485-501 (2000), Verma, Cancer Biochem. Biophys. 14: 151-162 (1994) and Dennis et al., Bioessays 5: 412-421 (1999).

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Another post-translational modification that may be altered in cancer cells is prenylation. Prenylation is the covalent attachment of a hydrophobic prenyl group (either farnesyl or geranylgeranyl) to a polypeptide. Prenylation is required for localizing a protein to a cell membrane and is often required for polypeptide function. For instance, the Ras superfamily of GTPase signalling proteins must be prenylated for function in a cell. See, e.g., Prendergast et al., Semin. Cancer Biol. 10: 443-452 (2000) and Khwaja et al., Lancet 355: 741-744 (2000).

Other post-translation modifications that may be altered in cancer cells include, without limitation, polypeptide methylation, acetylation, arginylation or racemization of amino acid residues. In these cases, the polypeptide from the cancerous cell may exhibit either increased or decreased amounts of the post-translational modification compared to the corresponding polypeptides from noncancerous cells.

Other polypeptide alterations in cancer cells include abnormal polypeptide cleavage of proteins and aberrant protein-protein interactions. Abnormal polypeptide cleavage may be cleavage of a polypeptide in a cancerous cell that does not usually occur in a normal cell, or a lack of cleavage in a cancerous cell, wherein the polypeptide is cleaved in a normal cell. Aberrant protein-protein interactions may be either covalent cross-linking or non-covalent binding between proteins that do not normally bind to each other. Alternatively, in a cancerous cell, a protein may fail to bind to another protein to which it is bound in a noncancerous cell. Alterations in cleavage or in protein-protein

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interactions may be due to over- or underproduction of a polypeptide in a cancerous cell compared to that in a normal cell, or may be due to alterations in post-translational modifications (see above) of one or more proteins in the cancerous cell. See, e.g., Henschen-Edman, Ann. N.Y. Acad. Sci. 936: 580-593 (2001).

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Alterations in polypeptide post-translational modifications, as well as changes in polypeptide cleavage and protein-protein interactions, may be determined by any method known in the art. For instance, alterations in phosphorylation may be determined by using anti-phosphoserine, anti-phosphothreonine or anti-phosphotyrosine antibodies or by amino acid analysis. Glycosylation alterations may be determined using antibodies specific for 10 different sugar residues, by carbohydrate sequencing, or by alterations in the size of the glycoprotein, which can be determined by, e.g., SDS polyacrylamide gel electrophoresis (PAGE). Other alterations of post-translational modifications, such as prenylation, racemization, methylation, acetylation and arginylation, may be determined by chemical analysis, protein sequencing, amino acid analysis, or by using antibodies specific for the particular post-translational modifications. Changes in protein-protein interactions and in polypeptide cleavage may be analyzed by any method known in the art including, without limitation, non-denaturing PAGE (for non-covalent protein-protein interactions), SDS PAGE (for covalent protein-protein interactions and protein cleavage), chemical cleavage. protein sequencing or immunoassays.

translationally modified. In one embodiment, polypeptides may be modified enzymatically or chemically, by addition or removal of a post-translational modification. For example, a polypeptide may be glycosylated or deglycosylated enzymatically. Similarly, polypeptides may be phosphorylated using a purified kinase, such as a MAP kinase (e.g., p38, ERK, or JNK) or a tyrosine kinase (e.g., Src or erbB2). A polypeptide may also be modified through synthetic chemistry. Alternatively, one may isolate the polypeptide of interest from a cell or tissue that expresses the polypeptide with the desired post-translational modification. In another embodiment, a nucleic acid molecule encoding the polypeptide of interest is introduced into a host cell that is capable of posttranslationally modifying the encoded polypeptide in the desired fashion. If the polypeptide does not contain a motif for a desired post-translational modification, one may alter the post-translational modification by mutating the nucleic acid sequence of a nucleic acid molecule encoding the polypeptide so that it contains a site for the desired post-

In another embodiment, the invention provides polypeptides that have been post-

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translational modification. Amino acid sequences that may be post-translationally modified are known in the art. See, e.g., the programs described above on the website www. expasy.org. The nucleic acid molecule may also be introduced into a host cell that is capable of post-translationally modifying the encoded polypeptide. Similarly, one may delete sites that are post-translationally modified by either mutating the nucleic acid sequence so that the encoded polypeptide does not contain the post-translational modification motif, or by introducing the native nucleic acid molecule into a host cell that is not capable of post-translationally modifying the encoded polypeptide.

It will be appreciated, as is well-known and as noted above, that polypeptides are not always entirely linear. For instance, polypeptides may be branched as a result of ubiquitination, and they may be circular, with or without branching, generally as a result of posttranslation events, including natural processing event and events brought about by human manipulation which do not occur naturally. Circular, branched and branched circular polypeptides may be synthesized by non-translation natural process and by entirely synthetic methods, as well. Modifications can occur anywhere in a polypeptide, including the peptide backbone, the amino acid side-chains and the amino or carboxyl termini. In fact, blockage of the amino or carboxyl group in a polypeptide, or both, by a covalent modification, is common in naturally occurring and synthetic polypeptides and such modifications may be present in polypeptides of the present invention, as well. For instance, the amino terminal residue of polypeptides made in *E. coli*, prior to proteolytic processing, almost invariably will be N-formylmethionine.

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Useful post-synthetic (and post-translational) modifications include conjugation to detectable labels, such as fluorophores. A wide variety of amine-reactive and thiol-reactive fluorophore derivatives have been synthesized that react under nondenaturing conditions with N-terminal amino groups and epsilon amino groups of lysine residues, on the one hand, and with free thiol groups of cysteine residues, on the other.

Kits are available commercially that permit conjugation of proteins to a variety of amine-reactive or thiol-reactive fluorophores: Molecular Probes, Inc. (Eugene, OR, USA), e.g., offers kits for conjugating proteins to Alexa Fluor 350, Alexa Fluor 430,

Fluorescein-EX, Alexa Fluor 488, Oregon Green 488, Alexa Fluor 532, Alexa Fluor 546, Alexa Fluor 546. Alexa Fluor 568. Alexa Fluor 594, and Texas Red-X.

A wide variety of other amine-reactive and thiol-reactive fluorophores are available commercially (Molecular Probes, Inc., Eugene, OR, USA), including Alexa 69

Fluor® 350, Alexa Fluor® 488, Alexa Fluor® 532, Alexa Fluor® 546, Alexa Fluor® 568, Alexa Fluor® 594, Alexa Fluor® 647 (monoclonal antibody labeling kits available from Molecular Probes, Inc., Eugene, OR, USA), BODIPY dyes, such as BODIPY 493/503, BODIPY FL, BODIPY R6G, BODIPY 530/550, BODIPY TMR, BODIPY 558/568,

BODIPY 558/568, BODIPY 564/570, BODIPY 576/589, BODIPY 581/591, BODIPY TR, BODIPY 630/650, BODIPY 650/665, Cascade Blue, Cascade Yellow, Dansyl, lissamine rhodamine B, Marina Blue, Oregon Green 488, Oregon Green 514, Pacific Blue, rhodamine 6G, rhodamine green, rhodamine red, tetramethylrhodamine, Texas Red (available from Molecular Probes, Inc., Eugene, OR, USA).

The polypeptides of the present invention can also be conjugated to fluorophores, other proteins, and other macromolecules, using bifunctional linking reagents. Common homobifunctional reagents include, e.g., APG, AEDP, BASED, BMB, BMDB, BMH, BMOE, BM[PEO]3, BM[PEO]4, BS3, BSOCOES, DFDNB, DMA, DMP, DMS, DPDPB, DSG, DSP (Lomant's Reagent), DSS, DST, DTBP, DTME, DTSSP, EGS, HBVS,

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Sulfo-BSOCOES, Sulfo-DST, Sulfo-EGS (all available from Pierce, Rockford, IL, USA); common heterobifunctional cross-linkers include ABH, AMAS, ANB-NOS, APDP, ASBA, BMPA, BMPH, BMPS, EDC, EMCA, EMCH, EMCS, KMUA, KMUH, GMBS, LC-SMCC, LC-SPDP, MBS, M2C2H, MPBH, MSA, NHS-ASA, PDPH, PMPI, SADP, SAED, SAND, SANPAH, SASD, SATP, SBAP, SFAD, SIA, SIAB, SMCC, SMPB,

20 SMPH, SMPT, SPDP, Sulfo-EMCS, Sulfo-GMBS, Sulfo-HSAB, Sulfo-KMUS, Sulfo-LC-SPDP, Sulfo-MBS, Sulfo-NHS-LC-ASA, Sulfo-SADP, Sulfo-SANPAH, Sulfo-SIAB, Sulfo-SMCC, Sulfo-SMPB, Sulfo-LC-SMPT, SVSB, TFCS (all available Pierce, Rockford, IL, USA).

Polypeptides of the present invention, including full length polypeptides, fragments and fusion proteins, can be conjugated, using such cross-linking reagents, to fluorophores that are not amine- or thiol-reactive. Other labels that usefully can be conjugated to polypeptides of the present invention include radioactive labels, echosonographic contrast reagents, and MRI contrast agents.

Polypeptides of the present invention, including full length polypeptide, fragments and fusion proteins, can also usefully be conjugated using cross-linking agents to carrier proteins, such as KLH, bovine thyroglobulin, and even bovine serum albumin (BSA), to increase immunogenicity for raising anti-CSP antibodies.

Polypeptides of the present invention, including full length polypeptide, fragments and fusion proteins, can also usefully be conjugated to polyethylene glycol (PEG): PEGylation increases the serum half life of proteins administered intravenously for replacement therapy. Delgado et al., Crit, Rev. Ther. Drug Carrier Syst. 9(3-4): 249-304 5 (1992); Scott et al., Curr. Pharm. Des. 4(6); 423-38 (1998); DeSantis et al., Curr. Opin. Biotechnol. 10(4): 324-30 (1999). PEG monomers can be attached to the protein directly or through a linker, with PEGylation using PEG monomers activated with tresyl chloride (2.2.2-trifluoroethanesulphonyl chloride) permitting direct attachment under mild conditions

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Polypeptides of the present invention are also inclusive of analogs of a polypeptide encoded by a nucleic acid molecule according to the instant invention. In a preferred embodiment, this polypeptide is a CSP. In a more preferred embodiment, this polypeptide is derived from a polypeptide having part or all of the amino acid sequence of SEO ID NO: 101-190. Also preferred is an analog polypeptide comprising one or more 15 substitutions of non-natural amino acids or non-native inter-residue bonds compared to the naturally-occurring polypeptide. In one embodiment, the analog is structurally similar to a CSP, but one or more peptide linkages is replaced by a linkage selected from the group consisting of --CH2NH--, --CH2S--, --CH2-CH2--, --CH=CH--(cis and trans), --COCH2--, --CH(OH)CH2-- and -CH2SO--. In another embodiment, the analog comprises substitution of one or more amino acids of a CSP with a D-amino acid of the same type or other non-natural amino acid in order to generate more stable peptides. D-amino acids can readily be incorporated during chemical peptide synthesis; peptides assembled from D-amino acids are more resistant to proteolytic attack; incorporation of D-amino acids can also be used to confer specific three dimensional conformations on the peptide. Other amino acid analogues commonly added during chemical synthesis include ornithine. norleucine, phosphorylated amino acids (typically phosphoserine, phosphothreonine, phosphotyrosine), L-malonyltyrosine, a non-hydrolyzable analog of phosphotyrosine (see, e.g., Kole et al., Biochem. Biophys. Res. Com. 209: 817-821 (1995)), and various halogenated phenylalanine derivatives.

Non-natural amino acids can be incorporated during solid phase chemical synthesis or by recombinant techniques, although the former is typically more common. Solid phase chemical synthesis of peptides is well established in the art. Procedures are described, inter alia, in Chan et al. (eds.), Fmoc Solid Phase Peptide Synthesis: A

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<u>Practical Approach</u> (Practical Approach Series), Oxford Univ. Press (March 2000); Jones, <u>Amino Acid and Peptide Synthesis</u> (Oxford Chemistry Primers, No 7), Oxford Univ. Press (1992); and Bodanszky, <u>Principles of Peptide Synthesis</u> (Springer Laboratory), Springer Verlag (1993).

Amino acid analogues having detectable labels are also usefully incorporated during synthesis to provide derivatives and analogs. Biotin, for example can be added using biotinoyl—(9-fluorenylmethoxycarbonyl)—Lysine (FMOC biocytin) (Molecular Probes, Eugene, OR, USA). Biotin can also be added enzymatically by incorporation into a fusion protein of a E. coli BirA substrate peptide. The FMOC and tBOC derivatives of dabcyl—Lysine (Molecular Probes, Inc., Eugene, OR, USA) can be used to incorporate the dabcyl chromophore at selected sites in the peptide sequence during synthesis. The aminonaphthalene derivative EDANS, the most common fluorophore for pairing with the dabcyl quencher in fluorescence resonance energy transfer (FRET) systems, can be introduced during automated synthesis of peptides by using EDANS—FMOC—L-glutamic acid or the corresponding tBOC derivative (both from Molecular Probes, Inc., Eugene, OR, USA). Tetramethylrhodamine fluorophores can be incorporated during automated FMOC synthesis of peptides using (FMOC)—TMR-L-lysine (Molecular Probes, Inc. Eugene, OR, USA).

Other useful amino acid analogues that can be incorporated during chemical synthesis include aspartic acid, glutamic acid, lysine, and tyrosine analogues having allyl side-chain protection (Applied Biosystems, Inc., Foster City, CA, USA); the allyl side chain permits synthesis of cyclic, branched-chain, sulfonated, glycosylated, and phosphorylated peptides.

A large number of other FMOC-protected non-natural amino acid analogues

25 capable of incorporation during chemical synthesis are available commercially, including,
e.g., Fmoc-2-aminobicyclo[2.2.1]heptane-2-carboxylic acid, Fmoc-3-exoaminobicyclo[2.2.1]heptane-2-endo-carboxylic acid, Fmoc-3-exoaminobicyclo[2.2.1]heptane-2-exo-carboxylic acid, Fmoc-3-exo-aminobicyclo[2.2.1]hept-5-ene-2-endo-carboxylic acid, Fmoc-3-exo-amino-bicyclo[2.2.1]hept5-ene-2-exo-carboxylic acid, Fmoc-cis-2-amino-1-cyclohexanecarboxylic acid, Fmoctrans-2-amino-1-cyclohexanecarboxylic acid, Fmoc-1-amino-1cyclopropanecarboxylic acid, Fmoc-D-2-amino-4-(ethylthiolbutyric acid, Fmoc-1-2-

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amino-4-(ethylthio)butyric acid, Fmoc-L-buthionine, Fmoc-S-methyl-L-Cysteine, Fmoc-2-aminobenzoic acid (anthranillic acid), Fmoc-3-aminobenzoic acid, Fmoc-4aminobenzoic acid. Fmoc-2-aminobenzophenone-2'-carboxylic acid. Fmoc-N-(4aminobenzovl)-B-alanine, Fmoc-2-amino-4.5-dimethoxybenzoic acid. Fmoc-4aminohippuric acid, Fmoc-2-amino-3-hydroxybenzoic acid, Fmoc-2-amino-5hydroxybenzoic acid, Fmoc-3-amino-4-hydroxybenzoic acid, Fmoc-4-amino-3hydroxybenzoic acid, Fmoc-4-amino-2-hydroxybenzoic acid, Fmoc-5-amino-2hydroxybenzoic acid, Fmoc-2-amino-3-methoxybenzoic acid, Fmoc-4-amino-3methoxybenzoic acid. Fmoc-2-amino-3-methylbenzoic acid. Fmoc-2-amino-5methylbenzoic acid, Fmoc-2-amino-6-methylbenzoic acid, Fmoc-3-amino-2-10 methylbenzoic acid, Fmoc-3-amino-4-methylbenzoic acid, Fmoc-4-amino-3methylbenzoic acid. Fmoc-3-amino-2-naphtoic acid. Fmoc-D.L-3-amino-3phenylpropionic acid, Fmoc-L-Methyldopa, Fmoc-2-amino-4,6-dimethyl-3pyridinecarboxylic acid, Fmoc-D,L-amino-2-thiophenacetic acid, Fmoc-4-15 (carboxymethyl)piperazine, Fmoc-4-carboxypiperazine, Fmoc-4-(carboxymethyl)homopiperazine, Fmoc-4-phenyl-4-piperidinecarboxylic acid, Fmoc-L-1.2.3.4-tetrahydronorharman-3-carboxylic acid. Fmoc-L-thiazolidine-4-carboxylic acid. all available from The Peptide Laboratory (Richmond, CA, USA).

Non-natural residues can also be added biosynthetically by engineering a suppressor tRNA, typically one that recognizes the UAG stop codon, by chemical aminoacylation with the desired unnatural amino acid. Conventional site-directed mutagenesis is used to introduce the chosen stop codon UAG at the site of interest in the protein gene. When the acylated suppressor tRNA and the mutant gene are combined in an in vitro transcription/translation system, the unnatural amino acid is incorporated in response to the UAG codon to give a protein containing that amino acid at the specified position. Liu et al., Proc. Natl Acad. Sci. USA 96(9): 4780-5 (1999); Wang et al., Science 292(5516): 498-500 (2001).

Fusion Proteins

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Another aspect of the present invention relates to the fusion of a polypeptide of the present invention to heterologous polypeptides. In a preferred embodiment, the polypeptide of the present invention is a CSP. In a more preferred embodiment, the polypeptide of the present invention that is fused to a heterologous polypeptide comprises part or all of the amino acid sequence of SEQ ID NO: 101-190, or is a mutein,

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homologous polypeptide, analog or derivative thereof. In an even more preferred embodiment, the fusion protein is encoded by a nucleic acid molecule comprising all or part of the nucleic acid sequence of SEQ ID NO: 1-100, or comprises all or part of a nucleic acid sequence that selectively hybridizes or is homologous to a nucleic acid molecule comprising a nucleic acid sequence of SEQ ID NO: 1-100.

The fusion proteins of the present invention will include at least one fragment of a polypeptide of the present invention, which fragment is at least 6, typically at least 8, often at least 15, and usefully at least 16, 17, 18, 19, or 20 amino acids long. The fragment of the polypeptide of the present to be included in the fusion can usefully be at least 25 amino acids long, at least 50 amino acids long, and can be at least 75, 100, or even 150 amino acids long. Fusions that include the entirety of a polypeptide of the present invention have particular utility.

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The heterologous polypeptide included within the fusion protein of the present invention is at least 6 amino acids in length, often at least 8 amino acids in length, and preferably at least 15, 20, or 25 amino acids in length. Fusions that include larger polypeptides, such as the IgG Fc region, and even entire proteins (such as GFP chromophore-containing proteins) are particularly useful.

As described above in the description of vectors and expression vectors of the present invention, which discussion is incorporated here by reference in its entirety, heterologous polypeptides to be included in the fusion proteins of the present invention can usefully include those designed to facilitate purification and/or visualization of recombinantly-expressed proteins. See, e.g., Ausubel, Chapter 16, (1992), supra. Although purification tags can also be incorporated into fusions that are chemically synthesized, chemical synthesis typically provides sufficient purity that further purification by HPLC suffices; however, visualization tags as above described retain their utility even when the protein is produced by chemical synthesis, and when so included render the fusion proteins of the present invention useful as directly detectable markers of the presence of a polypeptide of the invention.

As also discussed above, heterologous polypeptides to be included in the fusion proteins of the present invention can usefully include those that facilitate secretion of recombinantly expressed proteins into the periplasmic space or extracellular milieu for prokaryotic hosts or into the culture medium for eukaryotic cells through incorporation of secretion signals and/or leader sequences. For example, a His⁶ tagged protein can be

purified on a Ni affinity column and a GST fusion protein can be purified on a glutathione affinity column. Similarly, a fusion protein comprising the Fc domain of IgG can be purified on a Protein A or Protein G column and a fusion protein comprising an epitope tag such as myc can be purified using an immunoaffinity column containing an anti-c-myc antibody. It is preferable that the epitope tag be separated from the protein encoded by the essential gene by an enzymatic cleavage site that can be cleaved after purification. See also the discussion of nucleic acid molecules encoding fusion proteins that may be expressed on the surface of a cell.

Other useful fusion proteins of the present invention include those that permit use 10 of the polypeptide of the present invention as bait in a yeast two-hybrid system. See Bartel et al. (eds.), The Yeast Two-Hybrid System, Oxford University Press (1997): Zhu et al., Yeast Hybrid Technologies, Eaton Publishing (2000); Fields et al., Trends Genet. 10(8): 286-92 (1994); Mendelsohn et al., Curr. Opin. Biotechnol. 5(5): 482-6 (1994); Luban et al., Curr. Opin. Biotechnol. 6(1): 59-64 (1995); Allen et al., Trends Biochem. Sci. 20(12): 511-6 (1995); Drees, Curr. Opin. Chem. Biol. 3(1): 64-70 (1999); Topcu et al., Pharm. Res. 17(9): 1049-55 (2000); Fashena et al., Gene 250(1-2): 1-14 (2000); ; Colas et al., (1996) Genetic selection of peptide aptamers that recognize and inhibit cyclin-dependent kinase 2. Nature 380, 548-550; Norman, T. et al., (1999) Genetic selection of peptide inhibitors of biological pathways. Science 285, 591-595, Fabbrizio et al., (1999) Inhibition of mammalian cell proliferation by genetically selected peptide aptamers that functionally antagonize E2F activity. Oncogene 18, 4357-4363; Xu et al., (1997) Cells that register logical relationships among proteins. Proc Natl Acad Sci U S A. 94, 12473-12478; Yang, et al., (1995) Protein-peptide interactions analyzed with the yeast two-hybrid system. Nuc. Acids Res. 23, 1152-1156; Kolonin et al., (1998) Targeting 25 cyclin-dependent kinases in Drosophila with peptide aptamers. Proc Natl Acad Sci U S A 95, 14266-14271; Cohen et al., (1998) An artificial cell-cycle inhibitor isolated from a combinatorial library. Proc Natl Acad Sci USA 95, 14272-14277; Uetz, P.; Giot, L.; al, e.; Fields, S.; Rothberg, J. M. (2000) A comprehensive analysis of protein-protein interactions in Saccharomyces cerevisiae. Nature 403, 623-627; Ito, et al., (2001) A comprehensive two-hybrid analysis to explore the yeast protein interactome. Proc Natl Acad Sci USA 98, 4569-4574. Typically, such fusion is to either E. coli LexA or yeast GAL4 DNA binding domains. Related bait plasmids are available that express the bait fused to a nuclear localization signal.

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Other useful fusion proteins include those that permit display of the encoded polypeptide on the surface of a phage or cell, fusions to intrinsically fluorescent proteins, such as green fluorescent protein (GFP), and fusions to the IgG Fc region, as described above, which discussion is incorporated here by reference in its entirety.

The polypeptides of the present invention can also usefully be fused to protein toxins, such as Pseudomonas exotoxin A, diphtheria toxin, shiga toxin A, anthrax toxin lethal factor, ricin, in order to effect ablation of cells that bind or take up the proteins of the present invention.

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Fusion partners include, inter alia, myc, hemagglutinin (HA), GST, immunoglobulins, β -galactosidase, biotin trpE, protein Λ , β -lactamase, α -amylase, maltose binding protein, alcohol dehydrogenase, polyhistidine (for example, six histidine at the amino and/or carboxyl terminus of the polypeptide), lacZ, green fluorescent protein (GFP), yeast α mating factor, GAL4 transcription activation or DNA binding domain, luciferase, and serum proteins such as ovalbumin, albumin and the constant domain of IgG. See, e.g., Ausubel (1992), supra and Ausubel (1999), supra. Fusion proteins may also contain sites for specific enzymatic cleavage, such as a site that is recognized by enzymes such as a Factor XIII, trypsin, pepsin, or any other enzyme known in the art. Fusion proteins will typically be made by either recombinant nucleic acid methods, as described above, chemically synthesized using techniques well-known in the art (e.g., a Merrifield synthesis), or produced by chemical cross-linking.

Another advantage of fusion proteins is that the epitope tag can be used to bind the fusion protein to a plate or column through an affinity linkage for screening binding proteins or other molecules that bind to the CSP.

As further described below, the polypeptides of the present invention can readily be used as specific immunogens to raise antibodies that specifically recognize polypeptides of the present invention including CSPs and their allelic variants and homologues. The antibodies, in turn, can be used, inter alia, specifically to assay for the polypeptides of the present invention, particularly CSPs, e.g. by ELISA for detection of protein fluid samples, such as serum, by immunohistochemistry or laser scanning cytometry, for detection of protein in tissue samples, or by flow cytometry, for detection of intracellular protein in cell suspensions, for specific antibody-mediated isolation and/or purification of CSPs, as for example by immunoprecipitation, and for use as specific agonists or antagonists of CSPs.

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One may determine whether polypeptides of the present invention including CSPs, muteins, homologous proteins or allelic variants or fusion proteins of the present invention are functional by methods known in the art. For instance, residues that are tolerant of change while retaining function can be identified by altering the polypeptide at known residues using methods known in the art, such as alanine scanning mutagenesis, Cunningham et al., Science 244(4908): 1081-5 (1989); transposon linker scanning mutagenesis, Chen et al., Gene 263(1-2): 39-48 (2001); combinations of homolog- and alanine-scanning mutagenesis, Jin et al., J. Mol. Biol. 226(3): 851-65 (1992); combinatorial alanine scanning, Weiss et al., Proc. Natl. Acad. Sci USA 97(16): 8950-4 (2000), followed by functional assay. Transposon linker scanning kits are available commercially (New England Biolabs, Beverly, MA, USA, catalog. no. E7-102S; EZ::TNTM In-Frame Linker Insertion Kit, catalogue no. EZI04KN, Epicentre Technologies Corporation. Madison. WI, USA).

Purification of the polypeptides or fusion proteins of the present invention is well-known and within the skill of one having ordinary skill in the art. See, e.g., Scopes, Protein Purification, 2d ed. (1987). Purification of recombinantly expressed polypeptides is described above. Purification of chemically-synthesized peptides can readily be effected, e.g., by HPLC.

Accordingly, it is an aspect of the present invention to provide the isolated polypeptides or fusion proteins of the present invention in pure or substantially pure form in the presence of absence of a stabilizing agent. Stabilizing agents include both proteinaceous or non-proteinaceous material and are well-known in the art. Stabilizing agents, such as albumin and polyethylene glycol (PEG) are known and are commercially available.

Although high levels of purity are preferred when the isolated polypeptide or fusion protein of the present invention are used as therapeutic agents, such as in vaccines and replacement therapy, the isolated polypeptides of the present invention are also useful at lower purity. For example, partially purified polypeptides of the present invention can be used as immunogens to raise antibodies in laboratory animals.

In a preferred embodiment, the purified and substantially purified polypeptides of the present invention are in compositions that lack detectable ampholytes, acrylamide monomers, bis-acrylamide monomers, and polyacrylamide.

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The polypeptides or fusion proteins of the present invention can usefully be attached to a substrate. The substrate can be porous or solid, planar or non-planar; the bond can be covalent or noncovalent.

For example, the polypeptides or fusion proteins of the present invention can usefully be bound to a porous substrate, commonly a membrane, typically comprising nitrocellulose, polyvinylidene fluoride (PVDF), or cationically derivatized, hydrophilic PVDF; so bound, the polypeptides or fusion proteins of the present invention can be used to detect and quantify antibodies, e.g. in serum, that bind specifically to the immobilized polypeptide or fusion protein of the present invention.

As another example, the polypeptides or fusion proteins of the present invention can usefully be bound to a substantially nonporous substrate, such as plastic, to detect and quantify antibodies, e.g. in serum, that bind specifically to the immobilized protein of the present invention. Such plastics include polymethylacrylic, polyethylene, polypropylene, polyacrylate, polymethylmethacrylate, polyvinylchloride, polytetrafluoroethylene, polystyrene, polycarbonate, polyacetal, polysulfone, celluloscacetate, celluloscnitrate, nitrocellulose, or mixtures thereof, when the assay is performed in a standard microtiter dish, the plastic is typically polystyrene.

The polypeptides and fusion proteins of the present invention can also be attached to a substrate suitable for use as a surface enhanced laser desorption ionization source; so attached, the polypeptide or fusion protein of the present invention is useful for binding and then detecting secondary proteins that bind with sufficient affinity or avidity to the surface-bound polypeptide or fusion protein to indicate biologic interaction there between. The polypeptides or fusion proteins of the present invention can also be attached to a substrate suitable for use in surface plasmon resonance detection; so attached, the polypeptide or fusion protein of the present invention is useful for binding and then detecting secondary proteins that bind with sufficient affinity or avidity to the surface-bound polypeptide or fusion protein to indicate biological interaction there between.

Antibodies

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In another aspect, the invention provides antibodies, including fragments and derivatives thereof, that bind specifically to polypeptides encoded by the nucleic acid molecules of the invention. In a preferred embodiment, the antibodies are specific for a polypeptide that is a CSP, or a fragment, mutein, derivative, analog or fusion protein

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78 thereof. In a more preferred embodiment, the antibodies are specific for a polypeptide that comprises SEO ID NO: 101-190, or a fragment, mutein, derivative, analog or fusion protein thereof.

The antibodies of the present invention can be specific for linear epitopes, discontinuous epitopes, or conformational epitopes of such proteins or protein fragments. either as present on the protein in its native conformation or, in some cases, as present on the proteins as denatured, as, e.g., by solubilization in SDS. New epitopes may be also due to a difference in post translational modifications (PTMs) in disease versus normal tissue. For example, a particular site on a CSP may be glycosylated in cancerous cells, but not glycosylated in normal cells or vis versa. In addition, alternative splice forms of a CSP may be indicative of cancer. Differential degredation of the C or N-terminus of a CSP may also be a marker or target for anticancer therapy. For example, an CSP may be N-terminal degraded in cancer cells exposing new epitopes to which antibodies may selectively bind for diagnostic or therapeutic uses.

As is well-known in the art, the degree to which an antibody can discriminate as among molecular species in a mixture will depend, in part, upon the conformational relatedness of the species in the mixture; typically, the antibodies of the present invention will discriminate over adventitious binding to non-CSP polypeptides by at least two-fold. more typically by at least 5-fold, typically by more than 10-fold, 25-fold, 50-fold, 75-fold, and often by more than 100-fold, and on occasion by more than 500-fold or 1000-fold. When used to detect the proteins or protein fragments of the present invention, the antibody of the present invention is sufficiently specific when it can be used to determine the presence of the polypeptide of the present invention in samples derived from human colon.

Typically, the affinity or avidity of an antibody (or antibody multimer, as in the case of an IgM pentamer) of the present invention for a protein or protein fragment of the present invention will be at least about 1 x 10⁻⁶ molar (M), typically at least about 5 x 10⁻⁷ M, 1 x 10⁻⁷ M, with affinities and avidities of at least 1 x 10⁻⁸ M, 5 x 10⁻⁹ M, 1 x 10⁻¹⁰ M and up to 1 X 10⁻¹³ M proving especially useful.

The antibodies of the present invention can be naturally-occurring forms, such as IgG, IgM, IgD, IgE, IgY, and IgA, from any avian, reptilian, or mammalian species.

Human antibodies can, but will infrequently, be drawn directly from human donors or human cells. In such case, antibodies to the polypeptides of the present invention will

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typically have resulted from fortuitous immunization, such as autoimmune immunization, with the polypeptide of the present invention. Such antibodies will typically, but will not invariably, be polyclonal. In addition, individual polyclonal antibodies may be isolated and cloned to generate monoclonals.

Human antibodies are more frequently obtained using transgenic animals that express human immunoglobulin genes, which transgenic animals can be affirmatively immunized with the protein immunogen of the present invention. Human Ig-transgenic mice capable of producing human antibodies and methods of producing human antibodies therefrom upon specific immunization are described, *inter alia*, in United States Patent Nos. 6,162,963; 6,150,584; 6,114,598; 6,075,181; 5,939,598; 5,877,397; 5,874,299; 5,814,318; 5,789,650; 5,770,429; 5,661,016; 5,633,425; 5,625,126; 5,569,825; 5,545,807; 5,545,806, and 5,591,669, the disclosures of which are incorporated herein by reference in their entiretties. Such antibodies are typically monoclonal, and are typically produced using techniques developed for production of murine antibodies.

Human antibodies are particularly useful, and often preferred, when the antibodies of the present invention are to be administered to human beings as in vivo diagnostic or therapeutic agents, since recipient immune response to the administered antibody will often be substantially less than that occasioned by administration of an antibody derived from another species, such as mouse.

IgG, IgM, IgD, IgE, IgY, and IgA antibodies of the present invention are also usefully obtained from other species, including mammals such as rodents (typically mouse, but also rat, guinea pig, and hamster), lagomorphs,(typically rabbits), and also larger mammals, such as sheep, goats, cows, and horses; or egg laying birds or reptiles such as chickens or alligators. In such cases, as with the transgenic human-antibody-producing non-human mammals, fortuitous immunization is not required, and the non-human mammal is typically affirmatively immunized, according to standard immunization protocols, with the polypeptide of the present invention. One form of avian antibodies may be generated using techniques described in WO 00/29444, published 25 May 2000.

As discussed above, virtually all fragments of 8 or more contiguous amino acids of a polypeptide of the present invention can be used effectively as immunogens when conjugated to a carrier, typically a protein such as bovine thyroglobulin, keyhole limpet hemocyanin, or bovine serum albumin, conveniently using a bifunctional linker such as those described elsewhere above, which discussion is incorporated by reference here.

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Immunogenicity can also be conferred by fusion of the polypeptide of the present invention to other moieties. For example, polypeptides of the present invention can be produced by solid phase synthesis on a branched polylysine core matrix; these multiple antigenic peptides (MAPs) provide high purity, increased avidity, accurate chemical definition and improved safety in vaccine development. Tam et al., Proc. Natl. Acad. Sci. USA 85: 5409-5413 (1988): Posnett et al., J. Biol. Chem. 263: 1710-1725 (1988).

Protocols for immunizing non-human mammals or avian species are well-established in the art. See Harlow et al. (eds.), Using Antibodies: A Laboratory Manual, Cold Spring Harbor Laboratory (1998); Coligan et al. (eds.), Current Protocols in Immunology, John Wiley & Sons, Inc. (2001); Zola, Monoclonal Antibodies Preparation and Use of Monoclonal Antibodies and Engineered Antibody Derivatives (Basics: From Background to Bench), Springer Verlag (2000); Gross M, Speck J.Dtsch. Tierarztl. Wochenschr. 103: 417-422 (1996). Immunization protocols often include multiple immunizations, either with or without adjuvants such as Freund's complete adjuvant and Freund's incomplete adjuvant, and may include naked DNA immunization (Moss, Semin. Immunol. 2: 317-327 (1990).

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Antibodies from non-human mammals and avian species can be polyclonal or monoclonal, with polyclonal antibodies having certain advantages in immunohistochemical detection of the polypeptides of the present invention and monoclonal antibodies having advantages in identifying and distinguishing particular 20 epitopes of the polypeptides of the present invention. Antibodies from avian species may have particular advantage in detection of the polypeptides of the present invention, in human serum or tissues (Vikinge et al., Biosens. Bioelectron. 13: 1257-1262 (1998). Following immunization, the antibodies of the present invention can be obtained using any art-accepted technique. Such techniques are well-known in the art and are described in detail in references such as Coligan, supra; Zola, supra; Howard et al. (eds.), Basic Methods in Antibody Production and Characterization, CRC Press (2000); Harlow, supra; Davis (ed.), Monoclonal Antibody Protocols, Vol. 45, Humana Press (1995); Delves (ed.), Antibody Production: Essential Techniques, John Wiley & Son Ltd (1997); and Kenney. 30 Antibody Solution: An Antibody Methods Manual, Chapman & Hall (1997).

Briefly, such techniques include, *inter alia*, production of monoclonal antibodies by hybridomas and expression of antibodies or fragments or derivatives thereof from host cells engineered to express immunoglobulin genes or fragments thereof. These two methods of production are not mutually exclusive: genes encoding antibodies specific for the polypeptides of the present invention can be cloned from hybridomas and thereafter expressed in other host cells. Nor need the two necessarily be performed together: e.g., genes encoding antibodies specific for the polypeptides of the present invention can be cloned directly from B cells known to be specific for the desired protein, as further described in United States Patent No. 5,627,052, the disclosure of which is incorporated herein by reference in its entirety, or from antibody-displaying phage.

Recombinant expression in host cells is particularly useful when fragments or derivatives of the antibodies of the present invention are desired.

Host cells for recombinant antibody production of whole antibodies, antibody fragments, or antibody derivativescan be prokaryotic or eukaryotic.

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Prokaryotic hosts are particularly useful for producing phage displayed antibodies of the present invention.

The technology of phage-displayed antibodies, in which antibody variable region fragments are fused, for example, to the gene III protein (pIII) or gene VIII protein (pVIII) for display on the surface of filamentous phage, such as M13, is by now well-established. See, e.g., Sidhu, Curr. Opin. Biotechnol. 11(6): 610-6 (2000); Griffiths et al., Curr. Opin. Biotechnol. 9(1): 102-8 (1998); Hoogenboom et al., Immunotechnology, 4(1): 1-20 (1998); Rader et al., Current Opinion in Biotechnology 8: 503-508 (1997); Aujame et al., Human Antibodies 8: 155-168 (1997); Hoogenboom, Trends in Biotechnol. 15: 62-70 (1997); de Kruif et al., 17: 453-455 (1996); Barbas et al., Trends in Biotechnol. 14: 230-234 (1996); Winter et al., Ann. Rev. Immunol. 433-455 (1994). Techniques and protocols required to generate, propagate, screen (pan), and use the antibody fragments from such libraries have recently been compiled. See, e.g., Barbas (2001), supra; Kay, supra; and Abelson, supra.

Typically, phage-displayed antibody fragments are scFv fragments or Fab fragments; when desired, full length antibodies can be produced by cloning the variable regions from the displaying phage into a complete antibody and expressing the full length antibody in a further prokaryotic or a eukaryotic host cell. Eukaryotic cells are also useful for expression of the antibody fragments, and antibody derivatives of the present invention. For example, antibody fragments of the present invention can be produced in Pichia pastoris and in Saccharomyces cerevisiae. See, e.g., Takahashi et al., Biosci. Biotechnol. Biochem. 64(10): 2138-44 (2000); Freyre et al., J. Biotechnol. 76(2-3): 157-63 (2000); Fischer et al., Biotechnol. Appl. Biochem. 30 (Pt 2): 117-20

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(1999); Pennell et al., Res. Immunol. 149(6): 599-603 (1998); Eldin et al., J. Immunol. Methods. 201(1): 67-75 (1997);, Frenken et al., Res. Immunol. 149(6): 589-99 (1998); and Shusta et al., Nature Biotechnol. 16(8): 773-7 (1998).

Antibodies, including antibody fragments and derivatives, of the present invention can also be produced in insect cells. See, e.g., Li et al., Protein Expr. Purif. 21(1): 121-8 (2001); Ailor et al., Biotechnol. Bioeng. 58(2-3): 196-203 (1998); Hsu et al., Biotechnol. Prog. 13(1): 96-104 (1997); Edelman et al., Immunology 91(1): 13-9 (1997); and Nesbit et al., J. Immunol. Methods 151(1-2): 201-8 (1992).

Antibodies and fragments and derivatives thereof of the present invention can also be produced in plant cells, particularly maize or tobacco, Giddings et al., Nature Biotechnol. 18(11): 1151-5 (2000); Gavilondo et al., Biotechniques 29(1): 128-38 (2000); Fischer et al., J. Biol. Regul. Homeost. Agents 14(2): 83-92 (2000); Fischer et al., Biol. Chem. 30 (Pt 2): 113-6 (1999); Fischer et al., Biol. Chem. 380(7-8): 825-39 (1999); Russell, Curr. Top. Microbiol. Immunol. 240: 119-38 (1999); and Ma et al., Plant Physiol. 109(2): 341-6 (1995).

Antibodies, including antibody fragments and derivatives, of the present invention can also be produced in transgenic, non-human, mammalian milk. See, e.g. Pollock et al., J. Immunol Methods. 231: 147-57 (1999); Young et al., Res. Immunol. 149: 609-10 (1998); and Limonta et al., Immunotechnology 1: 107-13 (1995).

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Mammalian cells useful for recombinant expression of antibodies, antibody fragments, and antibody derivatives of the present invention include CHO cells, COS cells, 293 cells, and myeloma cells. Verma et al., J. Immunol. Methods 216(1-2):165-81 (1998) review and compare bacterial, yeast, insect and mammalian expression systems for expression of antibodies. Antibodies of the present invention can also be prepared by cell free translation, as further described in Merk et al., J. Biochem. (Tokyo) 125(2): 328-33 (1999) and Ryabova et al., Nature Biotechnol. 15(1): 79-84 (1997), and in the milk of transgenic animals, as further described in Pollock et al., J. Immunol. Methods 231(1-2): 147-57 (1999).

The invention further provides antibody fragments that bind specifically to one or more of the polypeptides of the present invention, to one or more of the polypeptides encoded by the isolated nucleic acid moleculess of the present invention, or the binding of which can be competitively inhibited by one or more of the polypeptides of the present invention or one or more of the polypeptides encoded by the isolated nucleic acid

molecules of the present invention. Among such useful fragments are Fab, Fab', Fv, F(ab)'2, and single chain Fv (soFv) fragments. Other useful fragments are described in Hudson. Curr. Opin. Biotechnol. 9(4): 395-402 (1998).

The present invention also relates to antibody derivatives that bind specifically to one or more of the polypeptides of the present invention, to one or more of the polypeptides encoded by the isolated nucleic acid molecules of the present invention, or the binding of which can be competitively inhibited by one or more of the polypeptides of the present invention or one or more of the polypeptides encoded by the isolated nucleic acid molecules of the present invention.

Among such useful derivatives are chimeric, primatized, and humanized antibodies; such derivatives are less immunogenic in human beings, and thus are more suitable for in vivo administration, than are unmodified antibodies from non-human mammalian species. Another useful derivative is PEGylation to increase the serum half life of the antibodies.

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15 Chimeric antibodies typically include heavy and/or light chain variable regions (including both CDR and framework residues) of immunoglobulins of one species, typically mouse, fused to constant regions of another species, typically human. See, e.g., Morrison et al., Proc. Natl. Acad. Sci USA.81(21): 6851-5 (1984); Sharon et al., Nature 309(5966); 364-7 (1984); Takeda et al., Nature 314(6010); 452-4 (1985); and United States Patent No. 5,807,715 the disclosure of which is incorporated herein by reference in its entirety. Primatized and humanized antibodies typically include heavy and/or light chain CDRs from a murine antibody grafted into a non-human primate or human antibody V region framework, usually further comprising a human constant region, Riechmann et al., Nature 332(6162): 323-7 (1988): Co et al., Nature 351(6326): 501-2 (1991): and United States Patent Nos. 6,054,297; 5,821,337; 5,770,196; 5,766,886; 5,821,123; 5,869,619; 6,180,377; 6,013,256; 5,693,761; and 6,180,370, the disclosures of which are incorporated herein by reference in their entireties. Other useful antibody derivatives of the invention include heteromeric antibody complexes and antibody fusions, such as diabodies (bispecific antibodies), single-chain diabodies, and intrabodies.

It is contemplated that the nucleic acids encoding the antibodies of the present invention can be operably joined to other nucleic acids forming a recombinant vector for cloning or for expression of the antibodies of the invention. Accordingly, the present invention includes any recombinant vector containing the coding sequences, or part

thereof, whether for eukaryotic transduction, transfection or gene therapy. Such vectors may be prepared using conventional molecular biology techniques, known to those with skill in the art, and would comprise DNA encoding sequences for the immunoglobulin V-regions including framework and CDRs or parts thereof, and a suitable promoter either with or without a signal sequence for intracellular transport. Such vectors may be transduced or transfected into eukaryotic cells or used for gene therapy (Marasco et al., Proc. Natl. Acad. Sci. (USA) 90: 7889-7893 (1993); Duan et al., Proc. Natl. Acad. Sci. (USA) 91: 5075-5079 (1994), by conventional techniques, known to those with skill in the art

The antibodies of the present invention, including fragments and derivatives thereof, can usefully be labeled. It is, therefore, another aspect of the present invention to provide labeled antibodies that bind specifically to one or more of the polypeptides of the present invention, to one or more of the polypeptides encoded by the isolated nucleic acid moleculess of the present invention, or the binding of which can be competitively inhibited by one or more of the polypeptides of the present invention or one or more of the polypeptides encoded by the isolated nucleic acid molecules of the present invention. The choice of label depends, in part, upon the desired use.

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For example, when the antibodies of the present invention are used for immunohistochemical staining of tissue samples, the label can usefully be an enzyme that catalyzes production and local deposition of a detectable product. Enzymes typically conjugated to antibodies to permit their immunohistochemical visualization are well-known, and include alkaline phosphatase, β-galactosidase, glucose oxidase, horseradish peroxidase (HRP), and urease. Typical substrates for production and deposition of visually detectable products include o-nitrophenyl-beta-D-galactopyranoside (ONPG); o-phenylenediamine dihydrochloride (OPD); p-nitrophenyl phosphate (PNPP); p-nitrophenyl-beta-D-galactopyranoside (PNPG); 3',3'-diaminobenzidine (DAB); 3-amino-9-ethylcarbazole (AEC); 4-chloro-1-naphthol (CN); 5-bromo-4-chloro-3-indolyl-phosphate (BCIP); ABTS®; BluoGal; iodonitrotetrazolium (INT); nitroblue tetrazolium chloride (NBT); phenazine methosulfate (PMS); phenolphthalein monophosphate (PMP); tetramethyl benzidine (TMB); tetranitroblue

tetrazolium (TNBT); X-Gal; X-Gluc; and X-Glucoside.

Other substrates can be used to produce products for local deposition that are luminescent. For example, in the presence of hydrogen peroxide (H₂O₂), horseradish

peroxidase (HRP) can catalyze the oxidation of cyclic diacylhydrazides, such as luminol. Immediately following the oxidation, the luminol is in an excited state (intermediate reaction product), which decays to the ground state by emitting light. Strong enhancement of the light emission is produced by enhancers, such as phenolic compounds. Advantages include high sensitivity, high resolution, and rapid detection without radioactivity and requiring only small amounts of antibody. See, e.g., Thorpe et al., Methods Enzymol. 133: 331-53 (1986); Kricka et al., J. Immunoassay 17(1): 67-83 (1996); and Lundqvist et al., J. Biolumin. Chemilumin. 10(6): 353-9 (1995). Kits for such enhanced chemiluminescent detection (ECL) are available commercially. The antibodies can also be labeled using 10 colloidal gold.

As another example, when the antibodies of the present invention are used, e.g., for flow cytometric detection, for seanning laser cytometric detection, or for fluorescent immunoassay, they can usefully be labeled with fluorophores. There are a wide variety of fluorophore labels that can usefully be attached to the antibodies of the present invention.

For flow cytometric applications, both for extracellular detection and for intracellular detection, common useful fluorophores can be fluorescein isothiocyanate (FITC), allophycocyanin (APC), R-phycocrythrin (PE), peridinin chlorophyll protein (PerCP),
Texas Red, Cy3, Cy5, fluorescence resonance energy tandem fluorophores such as PerCP-Cy5.5, PE-Cy5, PE-Cy5.5, PE-Cy7, PE-Texas Red, and APC-Cy7.

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Other fluorophores include, inter alia, Alexa Fluor® 350, Alexa Fluor® 488,
Alexa Fluor® 532, Alexa Fluor® 546, Alexa Fluor® 568, Alexa Fluor® 594, Alexa
Fluor® 647 (monoclonal antibody labeling kits available from Molecular Probes, Inc.,
Eugene, OR, USA), BODIPY dyes, such as BODIPY 493/503, BODIPY FL, BODIPY
R6G, BODIPY 530/550, BODIPY TMR, BODIPY 558/568, BODIPY 558/568, BODIPY
564/570, BODIPY 576/589, BODIPY 581/591, BODIPY TR, BODIPY 630/650,
BODIPY 650/665, Cascade Blue, Cascade Yellow, Dansyl, lissamine rhodamine B,
Marina Blue, Oregon Green 488, Oregon Green 514, Pacific Blue, rhodamine 6G,
rhodamine green, rhodamine red, tetramethylrhodamine, Texas Red (available from
Molecular Probes, Inc., Eugene, OR, USA), and Cy2, Cy3, Cy3.5, Cy5, Cy5.5, Cy7, all of
which are also useful for fluorescently labeling the antibodies of the present invention.
For secondary detection using labeled avidin, streptavidin, captavidin or neutravidin, the
antibodies of the present invention can usefully be labeled with biotin.

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When the antibodies of the present invention are used, e.g., for western blotting applications, they can usefully be labeled with radioisotopes, such as ³³P, ³²P, ³⁵S, ³H, and ¹²⁵I. As another example, when the antibodies of the present invention are used for radioimmunotherapy, the label can usefully be ²²⁸Th, ²²⁷Ac, ²²⁵Ac, ²²³Ra, ²¹³Bi, ²¹²Pb, ²¹²Bi, ²¹¹At, ²⁰³Pb, ¹⁹⁴OS, ¹⁸⁸Re, ¹⁸⁶Re, ¹³⁵Sm, ¹⁴⁹Tb, ¹³¹I, ¹²⁵I, ¹¹¹In, ¹⁰⁵Rh, ^{99m}Tc, ⁵⁷Ru, ⁹⁰Y, ⁹⁰Sr, ⁸⁸Y, ⁷²Se, ⁶⁷Cu, or ⁴⁷Sc.

As another example, when the antibodies of the present invention are to be used for in vivo diagnostic use, they can be rendered detectable by conjugation to MRI contrast agents, such as gadolinium diethylenetriaminepentaacetic acid (DTPA), Lauffer et al., Radiology 207(2): 529-38 (1998), or by radioisotopic labeling.

As would be understood, use of the labels described above is not restricted to the application as for which they were mentioned.

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The antibodies of the present invention, including fragments and derivatives thereof, can also be conjugated to toxins, in order to target the toxin's ablative action to cells that display and/or express the polypeptides of the present invention. Commonly, the antibody in such immunotoxins is conjugated to Pseudomonas exotoxin A, diphtheria toxin, shiga toxin A, anthrax toxin lethal factor, or ricin. See Hall (ed.), Immunotoxin Methods and Protocols (Methods in Molecular Biology, vol. 166), Humana Press (2000); and Frankel et al. (eds.), Clinical Applications of Immunotoxins. Springer-Verlag (1998).

The antibodies of the present invention can usefully be attached to a substrate, and it is, therefore, another aspect of the invention to provide antibodies that bind specifically to one or more of the polypeptides of the present invention, to one or more of the polypeptides encoded by the isolated nucleic acid molecules of the present invention, or the binding of which can be competitively inhibited by one or more of the polypeptides of the present invention or one or more of the polypeptides encoded by the isolated nucleic acid moleculess of the present invention, attached to a substrate. Substrates can be porous or nonporous, planar or nonplanar. For example, the antibodies of the present invention can usefully be conjugated to filtration media, such as NHS-activated Sepharose or CNBractivated Sepharose for purposes of immunoaffinity chromatography. For example, the antibodies of the present invention can usefully be attached to paramagnetic microspheres, typically by biotin-streptavidin interaction, which microsphere can then be used for isolation of cells that express or display the polypeptides of the present invention. As

another example, the antibodies of the present invention can usefully be attached to the surface of a microtiter plate for ELISA.

As noted above, the antibodies of the present invention can be produced in prokaryotic and eukaryotic cells. It is, therefore, another aspect of the present invention to provide cells that express the antibodies of the present invention, including hybridoma cells, B cells, plasma cells, and host cells recombinantly modified to express the antibodies of the present invention.

In yet a further aspect, the present invention provides aptamers evolved to bind specifically to one or more of the polypeptides of the present invention, to one or more of the polypeptides encoded by the isolated nucleic acid moleculess of the present invention, or the binding of which can be competitively inhibited by one or more of the polypeptides of the present invention or one or more of the polypeptides encoded by the isolated nucleic acid molecules of the present invention.

In sum, one of skill in the art, provided with the teachings of this invention, has available a variety of methods which may be used to alter the biological properties of the antibodies of this invention including methods which would increase or decrease the stability or half-life, immunogenicity, toxicity, affinity or yield of a given antibody molecule, or to alter it in any other way that may render it more suitable for a particular application.

20 Transgenic Animals and Cells

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In another aspect, the invention provides transgenic cells and non-human organisms comprising nucleic acid molecules of the invention. In a preferred embodiment, the transgenic cells and non-human organisms comprise a nucleic acid molecule encoding a CSP. In a preferred embodiment, the CSP comprises an amino acid sequence selected from SEQ ID NO: 101-190, or a fragment, mutein, homologous protein or allelic variant thereof. In another preferred embodiment, the transgenic cells and non-human organism comprise a CSNA of the invention, preferably a CSNA comprising a nucleotide sequence selected from the group consisting of SEQ ID NO: 1-100, or a part, substantially similar nucleic acid molecule, allelic variant or hybridizing nucleic acid molecule thereof.

In another embodiment, the transgenic cells and non-human organisms have a targeted disruption or replacement of the endogenous orthologue of the human CSG. The

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transgenic cells can be embryonic stem cells or somatic cells. The transgenic non-human organisms can be chimeric, nonchimeric heterozygotes, and nonchimeric homozygotes. Methods of producing transgenic animals are well-known in the art. See, e.g., Hogan et al., Manipulating the Mouse Embryo: A Laboratory Manual, 2d ed., Cold Spring Harbor Press (1999); Jackson et al., Mouse Genetics and Transgenics: A Practical Approach, Oxford University Press (2000); and Pinkert, Transgenic Animal Technology: A Laboratory Handbook, Academic Press (1999).

Any technique known in the art may be used to introduce a nucleic acid molecule of the invention into an animal to produce the founder lines of transgenic animals. Such techniques include, but are not limited to, pronuclear microinjection. (see, e.g., Paterson et al., Appl. Microbiol. Biotechnol. 40: 691-698 (1994); Carver et al., Biotechnology 11: 1263-1270 (1993); Wright et al., Biotechnology 9: 830-834 (1991); and United States Patent No. 4,873,191, herein incorporated by reference in its entirety); retrovirus-mediated gene transfer into germ lines, blastocysts or embryos (see, e.g., Van der Putten et al., Proc. Natl. Acad. Sci., USA 82: 6148-6152 (1985)); gene targeting in embryonic stem cells (see, e.g., Thompson et al., Cell 56: 313-321 (1989)); electroporation of cells or embryos (see, e.g., Lo, 1983, Mol. Cell. Biol. 3: 1803-1814 (1983)); introduction using a gene gun (see, e.g., Ulmer et al., Science 259: 1745-49 (1993); introducing nucleic acid constructs into embryonic pleuripotent stem cells and transferring the stem cells back into the blastocyst;

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Other techniques include, for example, nuclear transfer into enucleated oocytes of nuclei from cultured embryonic, fetal, or adult cells induced to quiescence (see, e.g., Campell et al., Nature 380: 64-66 (1996); Wilmut et al., Nature 385: 810-813 (1997)). The present invention provides for transgenic animals that carry the transgene (i.e., a nucleic acid molecule of the invention) in all their cells, as well as animals which carry the transcene in some, but not all their cells, i. e. mosaic animals or chimeric animals.

and sperm-mediated gene transfer (see, e.g., Lavitrano et al., Cell 57: 717-723 (1989)).

The transgene may be integrated as a single transgene or as multiple copies, such as in concatamers, e.g., head-to-head tandems or head-to-tail tandems. The transgene may also be selectively introduced into and activated in a particular cell type by following, e.g., the teaching of Lasko et al. et al., Proc. Natl. Acad. Sci. USA 89: 6232-6236 (1992). The regulatory sequences required for such a cell-type specific activation will depend upon the particular cell type of interest, and will be apparent to those of skill in the art.

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Once transgenic animals have been generated, the expression of the recombinant gene may be assayed utilizing standard techniques. Initial screening may be accomplished by Southern blot analysis or PCR techniques to analyze animal tissues to verify that integration of the transgene has taken place. The level of mRNA expression of the transgene in the tissues of the transgenic animals may also be assessed using techniques which include, but are not limited to, Northern blot analysis of tissue samples obtained from the animal, in situ hybridization analysis, and reverse transcriptase-PCR (RT-PCR). Samples of transgenic gene-expressing tissue may also be evaluated immunocytochemically or immunohistochemically using antibodies specific for the transgene product.

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Once the founder animals are produced, they may be bred, inbred, outbred, or crossbred to produce colonies of the particular animal. Examples of such breeding strategies include, but are not limited to: outbreeding of founder animals with more than one integration site in order to establish separate lines; inbreeding of separate lines in order to produce compound transgenics that express the transgene at higher levels because of the effects of additive expression of each transgene; crossing of heterozygous transgenic animals to produce animals homozygous for a given integration site in order to both augment expression and eliminate the need for screening of animals by DNA analysis; crossing of separate homozygous lines to produce compound heterozygous or homozygous lines; and breeding to place the transgene on a distinct background that is appropriate for an experimental model of interest.

Transgenic animals of the invention have uses which include, but are not limited to, animal model systems useful in elaborating the biological function of polypeptides of the present invention, studying conditions and/or disorders associated with aberrant expression, and in screening for compounds effective in ameliorating such conditions and/or disorders.

Methods for creating a transgenic animal with a disruption of a targeted gene are also well-known in the art. In general, a vector is designed to comprise some nucleotide sequences homologous to the endogenous targeted gene. The vector is introduced into a cell so that it may integrate, via homologous recombination with chromosomal sequences, into the endogenous gene, thereby disrupting the function of the endogenous gene. The transgene may also be selectively introduced into a particular cell type, thus inactivating the endogenous gene in only that cell type. See, e.g., Ou et al., Science 265: 103-106

(1994). The regulatory sequences required for such a cell-type specific inactivation will depend upon the particular cell type of interest, and will be apparent to those of skill in the art. See, e.g., Smithies et al., Nature 317: 230-234 (1985); Thomas et al., Cell 51: 503-512 (1987); Thompson et al., Cell 5: 313-321 (1989).

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In one embodiment, a mutant, non-functional nucleic acid molecule of the invention (or a completely unrelated DNA sequence) flanked by DNA homologous to the endogenous nucleic acid sequence (either the coding regions or regulatory regions of the gene) can be used, with or without a selectable marker and/or a negative selectable marker, to transfect cells that express polypeptides of the invention in vivo. In another embodiment, techniques known in the art are used to generate knockouts in cells that contain, but do not express the gene of interest. Insertion of the DNA construct, via targeted homologous recombination, results in inactivation of the targeted gene. Such approaches are particularly suited in research and agricultural fields where modifications to embryonic stem cells can be used to generate animal offspring with an inactive targeted gene. See, e.g., Thomas, supra and Thompson, supra. However this approach can be routinely adapted for use in humans provided the recombinant DNA constructs are directly administered or targeted to the required site in vivo using appropriate viral vectors that will be apparent to those of skill in the art.

In further embodiments of the invention, cells that are genetically engineered to express the polypeptides of the invention, or alternatively, that are genetically engineered not to express the polypeptides of the invention (e.g., knockouts) are administered to a patient in vivo. Such cells may be obtained from an animal or patient or an MHC compatible donor and can include, but are not limited to fibroblasts, bone marrow cells, blood cells (e.g., lymphocytes), adipocytes, muscle cells, endothelial cells etc. The cells are genetically engineered in vitro using recombinant DNA techniques to introduce the coding sequence of polypeptides of the invention into the cells, or alternatively, to disrupt the coding sequence and/or endogenous regulatory sequence associated with the polypeptides of the invention, e.g., by transduction (using viral vectors, and preferably vectors that integrate the transgene into the cell genome) or transfection procedures, including, but not limited to, the use of plasmids, cosmids, YACs, naked DNA, electroporation, liposomes, etc.

The coding sequence of the polypeptides of the invention can be placed under the control of a strong constitutive or inducible promoter or promoter/enhancer to achieve expression, and preferably secretion, of the polypeptides of the invention. The engineered cells which express and preferably secrete the polypeptides of the invention can be introduced into the patient systemically, e.g., in the circulation, or intraperitoneally.

Alternatively, the cells can be incorporated into a matrix and implanted in the body, e.g., genetically engineered fibroblasts can be implanted as part of a skin graft; genetically engineered endothelial cells can be implanted as part of a lymphatic or vascular graft. See, e.g., United States Patent Nos. 5,399,349 and 5,460,959, each of which is incorporated by reference herein in its entirety.

When the cells to be administered are non-autologous or non-MHC compatible cells, they can be administered using well-known techniques which prevent the development of a host immune response against the introduced cells. For example, the cells may be introduced in an encapsulated form which, while allowing for an exchange of components with the immediate extracellular environment, does not allow the introduced cells to be recognized by the host immune system.

Transgenic and "knock-out" animals of the invention have uses which include, but are not limited to, animal model systems useful in elaborating the biological function of polypeptides of the present invention, studying conditions and/or disorders associated with aberrant expression, and in screening for compounds effective in ameliorating such conditions and/or disorders.

20 Computer Readable Means

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A further aspect of the invention is a computer readable means for storing the nucleic acid and amino acid sequences of the instant invention. In a preferred embodiment, the invention provides a computer readable means for storing SEQ ID NO: 101-190 and SEQ ID NO: 1-100 as described herein, as the complete set of sequences or in any combination. The records of the computer readable means can be accessed for reading and display and for interface with a computer system for the application of programs allowing for the location of data upon a query for data meeting certain criteria, the comparison of sequences, the alignment or ordering of sequences meeting a set of criteria, and the like.

The nucleic acid and amino acid sequences of the invention are particularly useful as components in databases useful for search analyses as well as in sequence analysis algorithms. As used herein, the terms "nucleic acid sequences of the invention" and "amino acid sequences of the invention" mean any detectable chemical or physical characteristic of a polynucleotide or polypeptide of the invention that is or may be reduced to or stored in a computer readable form. These include, without limitation, chromatographic scan data or peak data, photographic data or scan data therefrom, and mass spectrographic data.

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This invention provides computer readable media having stored thereon sequences of the invention. A computer readable medium may comprise one or more of the following: a nucleic acid sequence comprising a sequence of a nucleic acid sequence of the invention; an amino acid sequence comprising an amino acid sequence of the invention; a set of nucleic acid sequences wherein at least one of said sequences comprises the sequence of a nucleic acid sequence of the invention; a set of amino acid sequences wherein at least one of said sequences comprises the sequence of an amino acid sequence of the invention; a data set representing a nucleic acid sequence comprising the sequence of one or more nucleic acid sequences of the invention; a data set representing a nucleic acid sequence encoding an amino acid sequence comprising the sequence of an amino acid sequence of the invention; a set of nucleic acid sequences wherein at least one of said sequences comprises the sequence of a nucleic acid sequence of the invention; a set of amino acid sequences wherein at least one of said sequences comprises the sequence of an amino acid sequence of the invention; a data set representing a nucleic acid sequence comprising the sequence of a nucleic acid sequence of the invention; a data set representing a nucleic acid sequence encoding an amino acid sequence comprising the sequence of an amino acid sequence of the invention. The computer readable medium can be any composition of matter used to store information or data, including, for example, commercially available floppy disks, tapes, hard drives, compact disks, and video disks.

Also provided by the invention are methods for the analysis of character sequences, particularly genetic sequences. Preferred methods of sequence analysis include, for example, methods of sequence bonology analysis, such as identity and similarity analysis, RNA structure analysis, sequence assembly, cladistic analysis, sequence motif analysis, open reading frame determination, nucleic acid base calling, and sequencing chromatogram peak analysis.

A computer-based method is provided for performing nucleic acid sequence identity or similarity identification. This method comprises the steps of providing a nucleic acid sequence comprising the sequence of a nucleic acid of the invention in a

computer readable medium; and comparing said nucleic acid sequence to at least one nucleic acid or amino acid sequence to identify sequence identity or similarity.

A computer-based method is also provided for performing amino acid homology identification, said method comprising the steps of: providing an amino acid sequence comprising the sequence of an amino acid of the invention in a computer readable medium; and comparing said amino acid sequence to at least one nucleic acid or an amino acid sequence to identify homology.

A computer-based method is still further provided for assembly of overlapping nucleic acid sequences into a single nucleic acid sequence, said method comprising the steps of: providing a first nucleic acid sequence comprising the sequence of a nucleic acid of the invention in a computer readable medium; and screening for at least one overlapping region between said first nucleic acid sequence and a second nucleic acid sequence. In addition, the invention includes a method of using patterns of expression associated with either the nucleic acids or proteins in a computer-based method to diagnose disease.

Diagnostic Methods for Colon Cancer

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The present invention also relates to quantitative and qualitative diagnostic assays and methods for detecting, diagnosing, monitoring, staging and predicting cancers by comparing expression of a CSNA or a CSP in a human patient that has or may have colon cancer, or who is at risk of developing colon cancer, with the expression of a CSNA or a CSP in a normal human control. For purposes of the present invention, "expression of a CSNA" or "CSNA expression" means the quantity of CSG mRNA that can be measured by any method known in the art or the level of transcription that can be measured by any 25 method known in the art in a cell, tissue, organ or whole patient. Similarly, the term "expression of a CSP" or "CSP expression" means the amount of CSP that can be measured by any method known in the art or the level of translation of a CSNA that can be measured by any method known in the art.

The present invention provides methods for diagnosing colon cancer in a patient. in particular squamous cell carcinoma, by analyzing for changes in levels of CSNA or CSP in cells, tissues, organs or bodily fluids compared with levels of CSNA or CSP in cells, tissues, organs or bodily fluids of preferably the same type from a normal human control, wherein an increase, or decrease in certain cases, in levels of a CSNA or CSP in

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the patient versus the normal human control is associated with the presence of colon cancer or with a predilection to the disease. In another preferred embodiment, the present invention provides methods for diagnosing colon cancer in a patient by analyzing changes in the structure of the mRNA of an CSG compared to the mRNA from a normal control. These changes include, without limitation, aberrant splicing, alterations in polyadenylation and/or alterations in 5' nucleotide capping. In yet another preferred embodiment, the present invention provides methods for diagnosing colon cancer in a patient by analyzing changes in a CSP compared to a CSP from a normal nationt. These changes include, e.g., alterations, including post translational modifications such as glycosylation and/or 10 phosphorylation of the CSP or changes in the subcellular CSP localization. For purposes of the present invention, diagnosing means that CSNA or CSP levels are used to determine the presence or absense of disease in a patient. As will be understood by those of skill in the art, measurement of other diagnostic parameters may be required for definitive diagnosis or determination of the appropriate treatment for the disease. The determination may be made by a clinician, a doctor, a testing laboratory, or a patient using an over the counter test. The patient may have symptoms of disease or may be assymptomatic. In addition, the CSNA or CSP levels of the present invention may be used as screening marker to determine whether further tests or biopsies are warranted. In addition, the CSNA or CSP levels may be used to determine the vunerability or

In a preferred embodiment, the expression of a CSNA is measured by determining the amount of a mRNA that encodes an amino acid sequence selected from SEQ ID NO: 101-190, a homolog, an allelic variant, or a fragment thereof. In a more preferred embodiment, the CSNA expression that is measured is the level of expression of a CSNA mRNA selected from SEQ ID NO: 1-100, or a hybridizing nucleic acid, homologous nucleic acid or allelic variant thereof, or a part of any of these nucleic acid, homologous nucleic acid or allelic variant thereof, or a part of any of these nucleic acid molecules. CSNA expression may be measured by any method known in the art, such as those described supra, including measuring mRNA expression by Northern blot, quantitative or qualitative reverse transcriptase PCR (RT-PCR), microarray, dot or slot blots or in situ hybridization. See, e.g., Ausubel (1992), supra; Ausubel (1999), supra; Sambrook (1989), supra; and Sambrook (2001), supra. CSNA transcription may be measured by any method known in the art including using a reporter gene hooked up to the promoter of a CSG of interest or doing nuclear run-off assavs. Alterations in mRNA structure, e.g.,

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susceptibility to disease.

aberrant splicing variants, may be determined by any method known in the art, including, RT-PCR followed by sequencing or restriction analysis. As necessary, CSNA expression may be compared to a known control, such as normal colon nucleic acid, to detect a change in expression.

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Id.

In another preferred embodiment, the expression of a CSP is measured by determining the level of a CSP having an amino acid sequence selected from the group consisting of SEQ ID NO: 101-190, a homolog, an allelic variant, or a fragment thereof. Such levels are preferably determined in at least one of cells, tissues, organs and/or bodily fluids, including determination of normal and abnormal levels. Thus, for instance, a diagnostic assay in accordance with the invention for diagnosing over- or underexpression of a CSNA or CSP compared to normal control bodily fluids, cells, or tissue samples may be used to diagnose the presence of colon cancer. The expression level of a CSP may be determined by any method known in the art, such as those described supra. In a preferred embodiment, the CSP expression level may be determined by radioimmunoassays. competitive-binding assays, ELISA, Western blot, FACS, immunohistochemistry, immunoprecipitation, proteomic approaches: two-dimensional gel electrophoresis (2D electrophoresis) and non-gel-based approaches such as mass spectrometry or protein interaction profiling. See, e.g, Harlow (1999), supra; Ausubel (1992), supra; and Ausubel (1999), supra. Alterations in the CSP structure may be determined by any method known in the art, including, e.g., using antibodies that specifically recognize phosphoserine, phosphothreonine or phosphotyrosine residues, two-dimensional polyacrylamide gel electrophoresis (2D PAGE) and/or chemical analysis of amino acid residues of the protein.

In a preferred embodiment, a radioimmunoassay (RIA) or an ELISA is used. An antibody specific to a CSP is prepared if one is not already available. In a preferred embodiment, the antibody is a monoclonal antibody. The anti-CSP antibody is bound to a solid support and any free protein binding sites on the solid support are blocked with a protein such as bovine serum albumin. A sample of interest is incubated with the antibody on the solid support under conditions in which the CSP will bind to the anti-CSP antibody. The sample is removed, the solid support is washed to remove unbound material, and an anti-CSP antibody that is linked to a detectable reagent (a radioactive substance for RIA and an enzyme for ELISA) is added to the solid support and incubated under conditions in which binding of the CSP to the labeled antibody will occur. After binding, the unbound

labeled antibody is removed by washing. For an ELISA, one or more substrates are added to produce a colored reaction product that is based upon the amount of an CSP in the sample. For an RIA, the solid support is counted for radioactive decay signals by any method known in the art. Quantitative results for both RIA and ELISA typically are obtained by reference to a standard curve.

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Other methods to measure CSP levels are known in the art. For instance, a competition assay may be employed wherein an anti-CSP antibody is attached to a solid support and an allocated amount of a labeled CSP and a sample of interest are incubated with the solid support. The amount of labeled CSP detected which is attached to the solid support can be correlated to the quantity of a CSP in the sample.

Of the proteomic approaches, 2D PAGE is a well-known technique. Isolation of individual proteins from a sample such as serum is accomplished using sequential separation of proteins by isoelectric point and molecular weight. Typically, polypeptides are first separated by isoelectric point (the first dimension) and then separated by size using an electric current (the second dimension). In general, the second dimension is perpendicular to the first dimension. Because no two proteins with different sequences are identical on the basis of both size and charge, the result of 2D PAGE is a roughly square gel in which each protein occupies a unique spot. Analysis of the spots with chemical or antibody probes, or subsequent protein microsequencing can reveal the relative abundance of a given protein and the identity of the proteins in the sample.

Expression levels of a CSNA can be determined by any method known in the art, including PCR and other nucleic acid methods, such as ligase chain reaction (LCR) and nucleic acid sequence based amplification (NASBA), can be used to detect malignant cells for diagnosis and monitoring of various malignancies. For example, reverse-transcriptase PCR (RT-PCR) is a powerful technique which can be used to detect the presence of a specific mRNA population in a complex mixture of thousands of other mRNA species. In RT-PCR, an mRNA species is first reverse transcribed to complementary DNA (cDNA) with use of the enzyme reverse transcriptase; the cDNA is then amplified as in a standard PCR reaction.

Hybridization to specific DNA molecules (e.g., oligonucleotides) arrayed on a solid support can be used to both detect the expression of and quantitate the level of expression of one or more CSNAs of interest. In this approach, all or a portion of one or more CSNAs is fixed to a substrate. A sample of interest, which may comprise RNA, e.g.,

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total RNA or polyA-selected mRNA, or a complementary DNA (cDNA) copy of the RNA is incubated with the solid support under conditions in which hybridization will occur between the DNA on the solid support and the nucleic acid molecules in the sample of interest. Hybridization between the substrate-bound DNA and the nucleic acid molecules in the sample can be detected and quantitated by several means, including, without limitation, radioactive labeling or fluorescent labeling of the nucleic acid molecule or a secondary molecule desirned to detect the hybrid.

The above tests can be carried out on samples derived from a variety of cells. bodily fluids and/or tissue extracts such as homogenates or solubilized tissue obtained from a patient. Tissue extracts are obtained routinely from tissue biopsy and autopsy material. Bodily fluids useful in the present invention include blood, urine, saliva or any other bodily secretion or derivative thereof. As used herein "blood" includes whole blood, plasma, serum or any derivative of blood. In a preferred embodiment, the specimen tested for expression of CSNA or CSP includes, without limitation, colon tissue, fluid obtained by bronchial alveolar lavage (BAL), sputum, colon cells grown in cell culture, blood, serum, lymph node tissue and lymphatic fluid. In another preferred embodiment, especially when metastasis of a primary colon cancer is known or suspected, specimens include, without limitation, tissues from brain, bone, bone marrow, liver, adrenal glands and colon. In general, the tissues may be sampled by biopsy, including, without limitation, needle biopsy, e.g., transthoracic needle aspiration, cervical mediatinoscopy, endoscopic lymph node biopsy, video-assisted thoracoscopy, exploratory thoracotomy, bone marrow biopsy and bone marrow aspiration. See Scott, supra and Franklin, pp. 529-570, in Kane, supra. For early and inexpensive detection, assaying for changes in CSNAs or CSPs in cells in sputum samples may be particularly useful. Methods of obtaining and analyzing sputum samples are disclosed in Franklin, supra.

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All the methods of the present invention may optionally include determining the expression levels of one or more other cancer markers in addition to determining the expression level of a CSNA or CSP. In many cases, the use of another cancer marker will decrease the likelihood of false positives or false negatives. In one embodiment, the one or more other cancer markers include other CSNA or CSPs as disclosed herein. Other cancer markers useful in the present invention will depend on the cancer being tested and are known to those of skill in the art. In a preferred embodiment, at least one other cancer marker in addition to a particular CSNA or CSP is measured. In a more preferred

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embodiment, at least two other additional cancer markers are used. In an even more preferred embodiment, at least three, more preferably at least five, even more preferably at least ten additional cancer markers are used.

Diagnosing

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In one aspect, the invention provides a method for determining the expression levels and/or structural alterations of one or more CSNA and/or CSP in a sample from a patient suspected of having colon cancer. In general, the method comprises the steps of obtaining the sample from the patient, determining the expression level or structural alterations of a CSNA and/or CSP and then ascertaining whether the patient has colon cancer from the expression level of the CSNA or CSP. In general, if high expression relative to a control of a CSNA or CSP is indicative of colon cancer, a diagnostic assay is considered positive if the level of expression of the CSNA or CSP is at least two times higher, and more preferably are at least five times higher, even more preferably at least ten times higher, than in preferably the same cells, tissues or bodily fluid of a normal human control. In contrast, if low expression relative to a control of a CSNA or CSP is indicative of colon cancer, a diagnostic assay is considered positive if the level of expression of the CSNA or CSP is at least two times lower, more preferably are at least five times lower, even more preferably at least ten times lower than in preferably the same cells, tissues or bodily fluid of a normal human control. The normal human control may be from a different patient or from uninvolved tissue of the same patient.

The present invention also provides a method of determining whether colon cancer has metastasized in a patient. One may identify whether the colon cancer has metastasized by measuring the expression levels and/or structural alterations of one or more CSNAs and/or CSPs in a variety of tissues. The presence of a CSNA or CSP in a certain tissue at levels higher than that of corresponding noncancerous tissue (e.g., the same tissue from another individual) is indicative of metastasis if high level expression of a CSNA or CSP is associated with colon cancer. Similarly, the presence of a CSNA or CSP in a tissue at levels lower than that of corresponding noncancerous tissue is indicative of metastasis if low level expression of a CSNA or CSP is associated with colon cancer. Further, the presence of a structurally altered CSNA or CSP that is associated with colon cancer is also indicative of metastasis.

In general, if high expression relative to a control of n CSNA or CSP is indicative of metastasis, an assay for metastasis is considered positive if the level of expression of

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the CSNA or CSP is at least two times higher, and more preferably are at least five times higher, even more preferably at least ten times higher, than in preferably the same cells, tissues or bodily fluid of a normal human control. In contrast, if low expression relative to a control of a CSNA or CSP is indicative of metastasis, an assay for metastasis is considered positive if the level of expression of the CSNA or CSP is at least two times lower, more preferably are at least five times lower, even more preferably at least ten times lower than in preferably the same cells, tissues or bodily fluid of a normal human control.

Staging

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The invention also provides a method of staging colon cancer in a human patient. The method comprises identifying a human patient having colon cancer and analyzing cells, tissues or bodily fluids from such human patient for expression levels and/or structural alterations of one or more CSNAs or CSPs. First, one or more tumors from a variety of patients are staged according to procedures well-known in the art, and the expression levels of one or more CSNAs or CSPs is determined for each stage to obtain a standard expression level for each CSNA and CSP. Then, the CSNA or CSP expression levels of the CSNA or CSP are determined in a biological sample from a patient whose stage of cancer is not known. The CSNA or CSP expression levels from the patient are then compared to the standard expression level. By comparing the expression level of the CSNAs and CSPs from the patient to the standard expression levels, one may determinethe stage of the tumor. The same procedure may be followed using structural alterations of a CSNA or CSP to determine the stage of a colon cancer.

Monitoring

Further provided is a method of monitoring colon cancer in a human patient. One may monitor a human patient to determine whether there has been metastasis and, if there has been, when metastasis began to occur. One may also monitor a human patient to determine whether a preneoplastic lesion has become cancerous. One may also monitor a human patient to determine whether a therapy, e.g., chemotherapy, radiotherapy or surgery, has decreased or eliminated the colon cancer. The monitoring may determine if there has been a reoccurrence and, if so, determine its nature. The method comprises identifying a human patient that one wants to monitor for colon cancer, periodically analyzing cells, tissues or bodily fluids from such human patient for expression levels of

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one or more CSNAs or CSPs, and comparing the CSNA or CSP levels over time to those CSNA or CSP expression levels obtained previously. Patients may also be monitored by measuring one or more structural alterations in a CSNA or CSP that are associated with colon cancer.

If increased expression of a CSNA or CSP is associated with metastasis, treatment failure, or conversion of a preneoplastic lesion to a cancerous lesion, then detecting an increase in the expression level of a CSNA or CSP indicates that the tumor is metastasizing, that treatment has failed or that the lesion is cancerous, respectively. One having ordinary skill in the art would recognize that if this were the case, then a decreased expression level would be indicative of no metastasis, effective therapy or failure to progress to a neoplastic lesion. If decreased expression of a CSNA or CSP is associated with metastasis, treatment failure, or conversion of a preneoplastic lesion to a cancerous lesion, then detecting an decrease in the expression level of a CSNA or CSP indicates that the tumor is metastasizing, that treatment has failed or that the lesion is cancerous, respectively. In a preferred embodiment, the levels of CSNAs or CSPs are determined from the same cell type, tissue or bodily fluid as prior patient samples. Monitoring a patient for onset of colon cancer metastasis is periodic and preferably is done on a quarterly basis, but may be done more or less frequently.

The methods described herein can further be utilized as prognostic assays to identify subjects having or at risk of developing a disease or disorder associated with increased or decreased expression levels of a CSNA and/or CSP. The present invention provides a method in which a test sample is obtained from a human patient and one or more CSNAs and/or CSPs are detected. The presence of higher (or lower) CSNA or CSP levels as compared to normal human controls is diagnostic for the human patient being at risk for developing cancer, particularly colon cancer. The effectiveness of therapeutic agents to decrease (or increase) expression or activity of one or more CSNAs and/or CSPs of the invention can also be monitored by analyzing levels of expression of the CSNAs and/or CSPs in a human patient in clinical trials or in in vitro screening assays such as in human cells. In this way, the gene expression pattern can serve as a marker, indicative of the physiological response of the human patient or cells, as the case may be, to the agent being tested.

Detection of Genetic Lesions or Mutations

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The methods of the present invention can also be used to detect genetic lesions or mutations in a CSG, thereby determining if a human with the genetic lesion is susceptible to developing colon cancer or to determine what genetic lesions are responsible, or are partly responsible, for a person's existing colon cancer. Genetic lesions can be detected, for example, by ascertaining the existence of a deletion, insertion and/or substitution of one or more nucleotides from the CSGs of this invention, a chromosomal rearrangement of a CSG, an aberrant modification of a CSG (such as of the methylation pattern of the genomic DNA), or allelic loss of a CSG. Methods to detect such lesions in the CSG of this invention are known to those having ordinary skill in the art following the teachings of the specification.

Methods of Detecting Noncancerous Colon Diseases

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The present invention also provides methods for determining the expression levels and/or structural alterations of one or more CSNAs and/or CSPs in a sample from a patient suspected of having or known to have a noncancerous colon disease. In general, the method comprises the steps of obtaining a sample from the patient, determining the expression level or structural alterations of a CSNA and/or CSP, comparing the expression level or structural alteration of the CSNA or CSP to a normal colon control, and then ascertaining whether the patient has a noncancerous colon disease. In general, if high expression relative to a control of a CSNA or CSP is indicative of a particular noncancerous colon disease, a diagnostic assay is considered positive if the level of expression of the CSNA or CSP is at least two times higher, and more preferably are at least five times higher, even more preferably at least ten times higher, than in preferably the same cells, tissues or bodily fluid of a normal human control. In contrast, if low expression relative to a control of a CSNA or CSP is indicative of a noncancerous colon disease, a diagnostic assay is considered positive if the level of expression of the CSNA or CSP is at least two times lower, more preferably are at least five times lower, even more preferably at least ten times lower than in preferably the same cells, tissues or bodily fluid of a normal human control. The normal human control may be from a different patient or from uninvolved tissue of the same patient.

One having ordinary skill in the art may determine whether a CSNA and/or CSP is associated with a particular noncancerous colon disease by obtaining colon tissue from a patient having a noncancerous colon disease of interest and determining which CSNAs

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and/or CSPs are expressed in the tissue at either a higher or a lower level than in normal colon tissue. In another embodiment, one may determine whether a CSNA or CSP exhibits structural alterations in a particular noncancerous colon disease state by obtaining colon tissue from a patient having a noncancerous colon disease of interest and determining the structural alterations in one or more CSNAs and/or CSPs relative to

Methods for Identifying Colon Tissue

normal colon tissue.

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In another aspect, the invention provides methods for identifying colon tissue.

These methods are particularly useful in, e.g., forensic science, colon cell differentiation and development, and in tissue engineering.

In one embodiment, the invention provides a method for determining whether a sample is colon tissue or has colon tissue-like characteristics. The method comprises the steps of providing a sample suspected of comprising colon tissue or having colon tissuelike characteristics, determining whether the sample expresses one or more CSNAs and/or CSPs, and, if the sample expresses one or more CSNAs and/or CSPs, concluding that the sample comprises colon tissue. In a preferred embodiment, the CSNA encodes a polypeptide having an amino acid sequence selected from SEO ID NO: 101-190, or a homolog, allelic variant or fragment thereof. In a more preferred embodiment, the CSNA has a nucleotide sequence selected from SEQ ID NO: 1-100, or a hybridizing nucleic acid, an allelic variant or a part thereof. Determining whether a sample expresses a CSNA can be accomplished by any method known in the art. Preferred methods include hybridization to microarrays. Northern blot hybridization, and quantitative or qualitative RT-PCR. In another preferred embodiment, the method can be practiced by determining 25 whether a CSP is expressed. Determining whether a sample expresses a CSP can be accomplished by any method known in the art. Preferred methods include Western blot. ELISA, RIA and 2D PAGE. In one embodiment, the CSP has an amino acid sequence selected from SEQ ID NO: 101-190, or a homolog, allelic variant or fragment thereof. In another preferred embodiment, the expression of at least two CSNAs and/or CSPs is 30 determined. In a more preferred embodiment, the expression of at least three, more preferably four and even more preferably five CSNAs and/or CSPs are determined.

In one embodiment, the method can be used to determine whether an unknown tissue is colon tissue. This is particularly useful in forensic science, in which small,

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damaged pieces of tissues that are not identifiable by microscopic or other means are recovered from a crime or accident scene. In another embodiment, the method can be used to determine whether a tissue is differentiating or developing into colon tissue. This is important in monitoring the effects of the addition of various agents to cell or tissue culture, e.g., in producing new colon tissue by tissue engineering. These agents include, e.g., growth and differentiation factors, extracellular matrix proteins and culture medium. Other factors that may be measured for effects on tissue development and differentiation include gene transfer into the cells or tissues, alterations in pH, aqueous:air interface and various other culture conditions.

10 Methods for Producing and Modifying Colon Tissue

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In another aspect, the invention provides methods for producing engineered colon tissue or cells. In one embodiment, the method comprises the steps of providing cells, introducing a CSNA or a CSG into the cells, and growing the cells under conditions in which they exhibit one or more properties of colon tissue cells. In a preferred embodiment, the cells are pluripotent. As is well-known in the art, normal colon tissue comprises a large number of different cell types. Thus, in one embodiment, the engineered colon tissue or cells comprises one of these cell types. In another embodiment, the engineered colon tissue or cells comprises more than one colon cell type. Further, the culture conditions of the cells or tissue may require manipulation in order to achieve full differentiation and development of the colon cell tissue. Methods for manipulating culture conditions are well-known in the art.

Nucleic acid molecules encoding one or more CSPs are introduced into cells, preferably pluripotent cells. In a preferred embodiment, the nucleic acid molecules encode CSPs having amino acid sequences selected from SEQ ID NO: 101-190, or homologous proteins, analogs, allelic variants or fragments thereof. In a more preferred embodiment, the nucleic acid molecules have a nucleotide sequence selected from SEQ ID NO: 1-100, or hybridizing nucleic acids, allelic variants or parts thereof. In another highly preferred embodiment, a CSG is introduced into the cells. Expression vectors and methods of introducing nucleic acid molecules into cells are well-known in the art and are described in detail. supra.

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Artificial colon tissue may be used to treat patients who have lost some or all of their colon function.

Pharmaceutical Compositions

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In another aspect, the invention provides pharmaceutical compositions comprising the nucleic acid molecules, polypeptides, fusion proteins, antibodies, antibody derivatives, antibody fragments, agonists, antagonists, or inhibitors of the present invention. In a preferred embodiment, the pharmaceutical composition comprises a CSNA or part thereof. In a more preferred embodiment, the CSNA has a nucleotide sequence selected from the group consisting of SEO ID NO: 1-100, a nucleic acid that hybridizes thereto, an allelic variant thereof, or a nucleic acid that has substantial sequence identity thereto. In another preferred embodiment, the pharmaceutical composition comprises a CSP or fragment thereof. In a more preferred embodiment, the pharmaceutical composition comprises a CSP having an amino acid sequence that is selected from the group consisting of SEO ID NO: 101-190, a polypeptide that is homologous thereto, a fusion protein comprising all or a portion of the polypeptide, or an analog or derivative thereof. In another preferred embodiment, the pharmaceutical composition comprises an anti-CSP antibody, preferably an antibody that specifically binds to a CSP having an amino acid that is selected from the group consisting of SEO ID NO: 101-190, or an antibody that binds to a polypeptide that is homologous thereto, a fusion protein comprising all or a portion of the polypeptide, or an analog or derivative thereof.

Such a composition typically contains from about 0.1 to 90% by weight of a therapeutic agent of the invention formulated in and/or with a pharmaceutically acceptable 25 carrier or excipient.

Pharmaceutical formulation is a well-established art that is further described in Gennaro (ed.), Remington: The Science and Practice of Pharmacy, 20th ed., Lippincott, Williams & Wilkins (2000); Ansel et al., Pharmaceutical Dosage Forms and Drug Delivery Systems, 7th ed., Lippincott Williams & Wilkins (1999); and Kibbe (ed.), Handbook of Pharmaceutical Excipients American Pharmaceutical Association, 3td ed. (2000) and thus need not be described in detail herein.

Briefly, formulation of the pharmaceutical compositions of the present invention will depend upon the route chosen for administration. The pharmaceutical compositions

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utilized in this invention can be administered by various routes including both enteral and parenteral routes, including oral, intravenous, intramuscular, subcutaneous, inhalation, topical, sublingual, rectal, intra-arterial, intramedullary, intrathecal, intraventricular, transmucosal, transdermal, intranasal, intraperitoneal, intrapulmonary, and intrauterine.

Oral dosage forms can be formulated as tablets, pills, dragees, capsules, liquids, gels, syrups, slurries, suspensions, and the like, for ingestion by the patient.

Solid formulations of the compositions for oral administration can contain suitable carriers or excipients, such as carbohydrate or protein fillers, such as sugars, including lactose, sucrose, mannitol, or sorbitol; starch from corn, wheat, rice, potato, or other plants; cellulose, such as methyl cellulose, hydroxypropylmethyl-cellulose, sodium carboxymethylcellulose, or microcrystalline cellulose; gums including arabic and tragacanth; proteins such as gelatin and collagen; inorganics, such as kaolin, calcium carbonate, dicalcium phosphate, sodium chloride; and other agents such as acacia and alginic acid.

Agents that facilitate disintegration and/or solubilization can be added, such as the cross-linked polyvinyl pyrrolidone, agar, alginic acid, or a salt thereof, such as sodium alginate, microcrystalline cellulose, corn starch, sodium starch glycolate, and alginic acid.

Tablet binders that can be used include acacia, methylcellulose, sodium carboxymethylcellulose, polyvinylpyrrolidone (PovidoneTM), hydroxypropyl methylcellulose, sucrose, starch and ethylcellulose.

Lubricants that can be used include magnesium stearates, stearic acid, silicone fluid, talc, waxes, oils, and colloidal silica.

Fillers, agents that facilitate disintegration and/or solubilization, tablet binders and lubricants, including the aforementioned, can be used singly or in combination.

Solid oral dosage forms need not be uniform throughout. For example, dragee cores can be used in conjunction with suitable coatings, such as concentrated sugar solutions, which can also contain gum arabic, tale, polyvinylpyrrolidone, carbopol gel, polyethylene glycol, and/or titanium dioxide, lacquer solutions, and suitable organic solvents or solvent mixtures.

Oral dosage forms of the present invention include push-fit capsules made of gelatin, as well as soft, sealed capsules made of gelatin and a coating, such as glycerol or sorbitol. Push-fit capsules can contain active ingredients mixed with a filler or binders, such as lactose or starches, lubricants, such as talc or magnesium stearate, and, optionally,

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stabilizers. In soft capsules, the active compounds can be dissolved or suspended in suitable liquids, such as fatty oils, liquid, or liquid polyethylene glycol with or without stabilizers.

Additionally, dyestuffs or pigments can be added to the tablets or dragec coatings for product identification or to characterize the quantity of active compound, i.e., dosage.

Liquid formulations of the pharmaceutical compositions for oral (enteral)
administration are prepared in water or other aqueous vehicles and can contain various
suspending agents such as methylcellulose, alginates, tragacanth, pectin, kelgin,
carrageenan, acacia, polyvinylpyrrolidone, and polyvinyl alcohol. The liquid formulations
oran also include solutions, emulsions, syrups and elixirs containing, together with the
active compound(s), wetting agents.

The pharmaceutical compositions of the present invention can also be formulated for parenteral administration. Formulations for parenteral administration can be in the form of aqueous or non-aqueous isotonic sterile injection solutions or suspensions.

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For intravenous injection, water soluble versions of the compounds of the present invention are formulated in, or if provided as a lyophilate, mixed with, a physiologically acceptable fluid vehicle, such as 5% dextrose ("D5"), physiologically buffered saline, 0.9% saline, Hanks' solution, or Ringer's solution. Intravenous formulations may include carriers, excipients or stabilizers including, without limitation, calcium, human serum albumin, citrate, acetate, calcium chloride, carbonate, and other salts.

Intramuscular preparations, e.g. a sterile formulation of a suitable soluble salt form of the compounds of the present invention, can be dissolved and administered in a pharmaceutical excipient such as Water-for-Injection, 0.9% saline, or 5% glucose solution. Alternatively, a suitable insoluble form of the compound can be prepared and administered as a suspension in an aqueous base or a pharmaceutically acceptable oil base, such as an ester of a long chain fatty acid (e.g., ethyl oleate), fatty oils such as sesame oil, triglycerides, or liposomes.

Parenteral formulations of the compositions can contain various carriers such as vegetable oils, dimethylacetamide, dimethylformamide, ethyl lactate, ethyl carbonate, isopropyl myristate, ethanol, polyols (glycerol, propylene glycol, liquid polyethylene glycol, and the like).

Aqueous injection suspensions can also contain substances that increase the viscosity of the suspension, such as sodium carboxymethyl cellulose, sorbitol, or dextran.

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Non-lipid polycationic amino polymers can also be used for delivery. Optionally, the suspension can also contain suitable stabilizers or agents that increase the solubility of the compounds to allow for the preparation of highly concentrated solutions.

Pharmaceutical compositions of the present invention can also be formulated to permit injectable, long-term, deposition. Injectable depot forms may be made by forming microencapsulated matrices of the compound in biodegradable polymers such as polylactide-polyglycolide. Depending upon the ratio of drug to polymer and the nature of the particular polymer employed, the rate of drug release can be controlled. Examples of other biodegradable polymers include poly(orthoesters) and poly(anhydrides). Depot injectable formulations are also prepared by entrapping the drug in microemulsions that are compatible with body tissues.

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The pharmaceutical compositions of the present invention can be administered topically. For topical use the compounds of the present invention can also be prepared in suitable forms to be applied to the skin, or mucus membranes of the nose and throat, and 15 can take the form of lotions, creams, ointments, liquid sprays or inhalants, drops, tinctures, lozenges. or throat paints. Such topical formulations further can include chemical compounds such as dimethylsulfoxide (DMSO) to facilitate surface penetration of the active ingredient. In other transdermal formulations, typically in patch-delivered formulations, the pharmaceutically active compound is formulated with one or more skin penetrants, such as 2-N-methyl-pyrrolidone (NMP) or Azone. A topical semi-solid ointment formulation typically contains a concentration of the active ingredient from about 1 to 20%, e.g., 5 to 10%, in a carrier such as a pharmaceutical cream base.

For application to the eyes or ears, the compounds of the present invention can be presented in liquid or semi-liquid form formulated in hydrophobic or hydrophilic bases as ointments, creams, lotions, paints or powders,

For rectal administration the compounds of the present invention can be administered in the form of suppositories admixed with conventional carriers such as cocoa butter, wax or other glyceride.

Inhalation formulations can also readily be formulated. For inhalation, various powder and liquid formulations can be prepared. For aerosol preparations, a sterile formulation of the compound or salt form of the compound may be used in inhalers, such as metered dose inhalers, and nebulizers. Aerosolized forms may be especially useful for treating respiratory disorders.

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Alternatively, the compounds of the present invention can be in powder form for reconstitution in the appropriate pharmaceutically acceptable carrier at the time of delivery.

The pharmaceutically active compound in the pharmaceutical compositions of the present invention can be provided as the salt of a variety of acids, including but not limited to hydrochloric, sulfuric, acetic, lactic, tartaric, malic, and succinic acid. Salts tend to be more soluble in aqueous or other protonic solvents than are the corresponding free base forms.

After pharmaceutical compositions have been prepared, they are packaged in an appropriate container and labeled for treatment of an indicated condition.

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The active compound will be present in an amount effective to achieve the intended purpose. The determination of an effective dose is well within the capability of those skilled in the art.

A "therapeutically effective dose" refers to that amount of active ingredient ,for

example CSP polypeptide, fusion protein, or fragments thereof, antibodies specific for

CSP, agonists, antagonists or inhibitors of CSP, which ameliorates the signs or symptoms

of the disease or prevent progression thereof; as would be understood in the medical arts,

cure, although desired, is not required.

The therapeutically effective dose of the pharmaceutical agents of the present invention can be estimated initially by in vitro tests, such as cell culture assays, followed by assay in model animals, usually mice, rats, rabbits, dogs, or pigs. The animal model can also be used to determine an initial preferred concentration range and route of administration.

For example, the ED50 (the dose therapeutically effective in 50% of the population) and LD50 (the dose lethal to 50% of the population) can be determined in one or more cell culture of animal model systems. The dose ratio of toxic to therapeutic effects is the therapeutic index, which can be expressed as LD50/ED50. Pharmaceutical compositions that exhibit large therapeutic indices are preferred.

The data obtained from cell culture assays and animal studies are used in formulating an initial dosage range for human use, and preferably provide a range of circulating concentrations that includes the ED50 with little or no toxicity. After administration, or between successive administrations, the circulating concentration of active agent varies within this range depending upon pharmacokinetic factors well-known

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in the art, such as the dosage form employed, sensitivity of the patient, and the route of administration.

The exact dosage will be determined by the practitioner, in light of factors specific to the subject requiring treatment. Factors that can be taken into account by the practitioner include the severity of the disease state, general health of the subject, age, weight, gender of the subject, diet, time and frequency of administration, drug combination(s), reaction sensitivities, and tolerance/response to therapy. Long-acting pharmaceutical compositions can be administered every 3 to 4 days, every week, or once every two weeks depending on half-life and clearance rate of the particular formulation.

Normal dosage amounts may vary from 0.1 to 100,000 micrograms, up to a total dose of about 1 g, depending upon the route of administration. Where the therapeutic agent is a protein or antibody of the present invention, the therapeutic protein or antibody agent typically is administered at a daily dosage of 0.01 mg to 30 mg/kg of body weight of the patient (e.g., 1mg/kg to 5 mg/kg). The pharmaceutical formulation can be 15 administered in multiple doses per day, if desired, to achieve the total desired daily dose.

Guidance as to particular dosages and methods of delivery is provided in the literature and generally available to practitioners in the art. Those skilled in the art will employ different formulations for nucleotides than for proteins or their inhibitors. Similarly, delivery of polynucleotides or polypeptides will be specific to particular cells, conditions, locations, etc.

Conventional methods, known to those of ordinary skill in the art of medicine, can be used to administer the pharmaceutical formulation(s) of the present invention to the patient. The pharmaceutical compositions of the present invention can be administered alone, or in combination with other therapeutic agents or interventions.

25 Therapeutic Methods

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The present invention further provides methods of treating subjects having defects in a gene of the invention, e.g., in expression, activity, distribution, localization, and/or solubility, which can manifest as a disorder of colon function. As used herein, "treating" includes all medically-acceptable types of therapeutic intervention, including palliation and prophylaxis (prevention) of disease. The term "treating" encompasses any improvement of a disease, including minor improvements. These methods are discussed below.

Gene Therapy and Vaccines

The isolated nucleic acids of the present invention can also be used to drive in vivo expression of the polypeptides of the present invention. In vivo expression can be driven from a vector, typically a viral vector, often a vector based upon a replication incompetent retrovirus, an adenovirus, or an adeno-associated virus (AAV), for the purpose of gene therapy. In vivo expression can also be driven from signals endogenous to the nucleic acid or from a vector, often a plasmid vector, such as pVAX1 (Invitrogen, Carlsbad, CA, USA), for purpose of "naked" nucleic acid vaccination, as further described in United States Patent Nos. 5,589,466; 5,679,647; 5,804,566; 5,830,877; 5,843,913; 5,880,104; 10 5.958.891; 5.985.847; 6.017.897; 6.110.898; 6.204.250, the disclosures of which are incorporated herein by reference in their entireties. For cancer therapy, it is preferred that the vector also be tumor-selective. See, e.g., Doronin et al., J. Virol, 75: 3314-24 (2001).

In another embodiment of the therapeutic methods of the present invention, a therapeutically effective amount of a pharmaceutical composition comprising a nucleic 15 acid molecule of the present invention is administered. The nucleic acid molecule can be delivered in a vector that drives expression of a CSP, fusion protein, or fragment thereof, or without such vector. Nucleic acid compositions that can drive expression of a CSP are administered, for example, to complement a deficiency in the native CSP, or as DNA vaccines. Expression vectors derived from virus, replication deficient retroviruses, adenovirus, adeno-associated (AAV) virus, herpes virus, or vaccinia virus can be used as can plasmids. See, e.g., Cid-Arregui, supra. In a preferred embodiment, the nucleic acid molecule encodes a CSP having the amino acid sequence of SEQ ID NO: 101-190, or a fragment, fusion protein, allelic variant or homolog thereof.

In still other therapeutic methods of the present invention, pharmaceutical compositions comprising host cells that express a CSP, fusions, or fragments thereof can be administered. In such cases, the cells are typically autologous, so as to circumvent xenogeneic or allotypic rejection, and are administered to complement defects in CSP production or activity. In a preferred embodiment, the nucleic acid molecules in the cells encode a CSP having the amino acid sequence of SEO ID NO: 101-190, or a fragment. fusion protein, allelic variant or homolog thereof.

Antisense Administration

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Antisense nucleic acid compositions, or vectors that drive expression of a CSG antisense nucleic acid, are administered to downregulate transcription and/or translation of

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a CSG in circumstances in which excessive production, or production of aberrant protein, is the pathophysiologic basis of disease.

Antisense compositions useful in therapy can have a sequence that is complementary to coding or to noncoding regions of a CSG. For example,

oligonucleotides derived from the transcription initiation site, e.g., between positions -10 and +10 from the start site, are preferred.

Catalytic antisense compositions, such as ribozymes, that are capable of sequence-specific hybridization to CSG transcripts, are also useful in therapy. See, e.g., Phylactou, Adv. Drug Deliv. Rev. 44(2-3): 97-108 (2000); Phylactou et al., Hum. Mol. Genet. 7(10): 1649-53 (1998); Rossi, Ciba Found. Symp. 209: 195-204 (1997); and Sigurdsson et al., Trends Biotechnol. 13(8): 286-9 (1995).

Other nucleic acids useful in the therapeutic methods of the present invention are those that are capable of triplex helix formation in or near the CSG genomic locus. Such triplexing oligonucleotides are able to inhibit transcription. See, e.g., Intody et al., Nucleic Acids Res. 28(21): 4283-90 (2000); and McGuffie et al., Cancer Res. 60(14): 3790-9 (2000). Pharmaceutical compositions comprising such triplex forming oligos (TFOs) are administered in circumstances in which excessive production, or production of aberrant protein, is a pathophysiologic basis of disease.

In a preferred embodiment, the antisense molecule is derived from a nucleic acid molecule encoding a CSP, preferably a CSP comprising an amino acid sequence of SEQ ID NO: 101-190, or a fragment, allelic variant or homolog thereof. In a more preferred embodiment, the antisense molecule is derived from a nucleic acid molecule having a nucleotide sequence of SEQ ID NO: 1-100, or a part, allelic variant, substantially similar or hybridizing nucleic acid thereof.

Polypeptide Administration

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In one embodiment of the therapeutic methods of the present invention, a therapeutically effective amount of a pharmaceutical composition comprising a CSP, a fusion protein, fragment, analog or derivative thereof is administered to a subject with a clinically-significant CSP defect.

Protein compositions are administered, for example, to complement a deficiency in native CSP. In other embodiments, protein compositions are administered as a vaccine to elicit a humoral and/or cellular immune response to CSP. The immune response can be used to modulate activity of CSP or, depending on the immunogen, to immunize against

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aberrant or aberrantly expressed forms, such as mutant or inappropriately expressed isoforms. In yet other embodiments, protein fusions having a toxic moiety are administered to ablate cells that aberrantly accumulate CSP.

In a preferred embodiment, the polypeptide administered is a CSP comprising an amino acid sequence of SEQ ID NO: 101-190, or a fusion protein, allelic variant, homolog, analog or derivative thereof. In a more preferred embodiment, the polypeptide is encoded by a nucleic acid molecule having a nucleide sequence of SEQ ID NO: 1-100, or a part, allelic variant, substantially similar or hybridizing nucleic acid thereof.

Antibody, Agonist and Antagonist Administration

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In another embodiment of the therapeutic methods of the present invention, a therapeutically effective amount of a pharmaceutical composition comprising an antibody (including fragment or derivative thereof) of the present invention is administered. As is well-known, antibody compositions are administered, for example, to antagonize activity of CSP, or to target therapeutic agents to sites of CSP presence and/or accumulation. In a preferred embodiment, the antibody specifically binds to a CSP comprising an amino acid sequence of SEQ ID NO: 101-190, or a fusion protein, allelic variant, homolog, analog or derivative thereof. In a more preferred embodiment, the antibody specifically binds to a CSP encoded by a nucleic acid molecule having a nucleotide sequence of SEQ ID NO: 1-100, or a part, allelic varian, substantially similar or hybridizing nucleic acid thereof.

The present invention also provides methods for identifying modulators which bind to a CSP or have a modulatory effect on the expression or activity of a CSP. Modulators which decrease the expression or activity of CSP (antagonists) are believed to be useful in treating colon cancer. Such screening assays are known to those of skill in the art and include, without limitation, cell-based assays and cell-free assays. Small molecules predicted via computer imaging to specifically bind to regions of a CSP can also be designed, synthesized and tested for use in the imaging and treatment of colon cancer. Further, libraries of molecules can be screened for potential anticancer agents by assessing the ability of the molecule to bind to the CSPs identified herein. Molecules identified in the library as being capable of binding to a CSP are key candidates for further evaluation for use in the treatment of colon cancer. In a preferred embodiment, these molecules will downregulate expression and/or activity of a CSP in cells.

In another embodiment of the therapeutic methods of the present invention, a pharmaceutical composition comprising a non-antibody antagonist of CSP is administered.

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Antagonists of CSP can be produced using methods generally known in the art. In particular, purified CSP can be used to screen libraries of pharmaceutical agents, often combinatorial libraries of small molecules, to identify those that specifically bind and antagonize at least one activity of a CSP.

In other embodiments a pharmaceutical composition comprising an agonist of a CSP is administered. Agonists can be identified using methods analogous to those used to identify antagonists.

In a preferred embodiment, the antagonist or agonist specifically binds to and antagonizes or agonizes, respectively, a CSP comprising an amino acid sequence of SEQ ID NO: 101-190, or a fusion protein, allelic variant, homolog, analog or derivative thereof. In a more preferred embodiment, the antagonist or agonist specifically binds to and antagonizes or agonizes, respectively, a CSP encoded by a nucleic acid molecule having a nucleotide sequence of SEQ ID NO: 1-100, or a part, allelic variant, substantially similar or hybridizing nucleic acid thereof.

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Targeting Colon Tissue

The invention also provides a method in which a polypeptide of the invention, or an antibody thereto, is linked to a therapeutic agent such that it can be delivered to the colon or to specific cells in the colon. In a preferred embodiment, an anti-CSP antibody is linked to a therapeutic agent and is administered to a patient in need of such therapeutic agent. The therapeutic agent may be a toxin, if colon tissue needs to be selectively destroyed. This would be useful for targeting and killing colon cancer cells. In another embodiment, the therapeutic agent may be a growth or differentiation factor, which would be useful for promoting colon cell function.

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In another embodiment, an anti-CSP antibody may be linked to an imaging agent that can be detected using, e.g., magnetic resonance imaging, CT or PET. This would be useful for determining and monitoring colon function, identifying colon cancer tumors, and identifying noncancerous colon diseases.

114 EXAMPLES

Example 1: Gene Expression analysis

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Identification of CSGs was carried out by a systematic analysis of gene expression data in the LIFESEQ® Gold database available from Incyte Genomics Inc, Palo Alto, CA, using the data mining software package CLASP™.

The CLASP target gene identification process is focused on, but not limited to, the following 5 CLASP profiles: tissue specific expression, cancer specific expression, differentially expressed in cancer, maximum tissue differential expression.

- (1) For these profiles:cDNA libraries were divided into 48 unique
 10 tissue organs. The genes were grouped into gene bins, each bin is a
 sequence based cluster grouped together with a common contig.
 The expression levels for each gene bin were calculated in each
 organ. Differential expression significance was calculated with
 rigorous statistical significant test considering the influence of
 sequence random fluctuations and sampling size of cDNA libraries
 from concept published by Audic S and Claverie JM (Genome Res
 1997 7(10): 986-995: The significance of digital gene expression
 profiles).
- (2) Highly expressed organ specific genes were selected based on the
 20 percentage abundance level in the targeted organ versus all the other
 organs (organ-specificity).
 - (3) The expression levels of each highly expressed organ-specific gene in the tumor tissue libraries were compared with normal tissue libraries and tissue libraries associated with tumor or disease (cancer-specificity) and analyzed for statistical significance.
 - (4) Target genes exhibiting each CLASP profile criteria were selected

CLASP 1 tissue specific expression profile: In order to meet the organ-specificity criteria, the expression level of the component clones which the gene is composed of must exhibit 3 or more occurrences regardless the total number of genes isolated for the target organ. The percentage abundance level in each organ was calculated to identify the organ with the highest expression percentage level.

CLASP 2 cancer specific expression profile: In order to fulfill the cancer specific criteria, genes must exhibit 0 expression in normal and libraries associated with tumor and disease but not tumor per se. If the gene then exhibited organ-specificity, the gene was selected as a CLASP target for this profile.

CLASP 3 maximum tissue differential expression profile: CLASP targets were selected based on ratio of expression in tumor libraries compared to expression in normal libraries (including normal libraries associated with tumor or disease) for each organ regardless of whether the gene exhibited organ-specificity. This profile was divided into 2 sub-profiles, since the ratio of expression cannot be obtained if no expression is present in normal libraries (including normal libraries associated with tumor or disease). In this case, the maximum expression percentage of the gene, as calculated by the occurrence of the gene divided by the occurrence of all genes in the target organ, was used. CLASP selects the ton 50 targets for each sub-profile.

CLASP 4 maximum tissue differential expression profile with negligible
expression in normal tissues: CLASP targets were selected based on ratio of expression in
tumor libraries compared to expression in normal libraries (including normal libraries

associated with tumor or disease) for each organ regardless of whether the gene exhibited
organ-specificity.

CLASP 5 differentially expressed in cancer profile: Expression levels in tumor libraries in each organ and normal libraries (including normal libraries associated with cancer or disease) for all organs were obtained and statistically analyzed. If the gene exhibited 90% of confidence that it is over-expressed in tumor libraries in the target organ than normal libraries for all organs, it was selected as a CLASP target for this profile. Accordingly, CLASP allows the identification of highly expressed organ and cancer specific genes based on the gene expression levels in each tissue organ. CLASP scores for a portion of the CSG of this invention are listed below.

25 The CLASP scores for SEQ ID NO: 1-100 are listed below:

SEQ ID	Organ	Abundance - Percentage	Organ	Abundance – Percentage	Organ	Abundance – Percentage	Organ	Abundance – Percentage
NO:	INS	30028						
NO: 2	INS	30028	FTS	20003				
NO:	INS	30028	SKN	2003	EAE	10062		
NO:	INS	30028	UTR	10004	OVR	10007		
NO: 5	INS	30028	INL	10004	LNG	10003	TST	10011
NO:	INS	30028	BRN	10001	THR	1002		
NO:	INS	30028						
NO:	INS	20257						

8							
NO:	INL	5002	INS	30028	THY	10019	
NO: 10	INS	50047	INL	10004			
NO: 11	INL	5002	TNS	30049	UNC	10011	
NO: 12	INL	40016					
NO: 13	INL	30012					
NO: 14	INL	40016					
NO: 15	INS	50047	INL	5002	PAN	10008	
NO: 16	INS	50047	INL	5002			
NO: 17	INL	30012					
NO: 18	INL	30012					
NO: 19	INL	30012	222				
NO: 20	INL	30012	PRO	10003			
NO: 21	INL	30012					
NO: 22	INL	20043					
NO: 23 NO:	INS	30028 80032			_		
24	INL						
NO: 25		80032					
NO: 26	INL	80032					
27 NO:	INL	30012					
28 NO:	INL	20047					
29 NO:	INL	90036	INS	10009	GEM	20042	
30 NO:	INL	90036	INS	10009	GEN	20042	
31 NO:	INL	20059	1110	10009			
32 NO:	INL	30012					
33 NO:	INS	30028	MSL	1002	BLV	10006	
34 NO:	INS	30028		1 - 1002		1 .0000	
35 NO:	INS	20162					
36 NO:	INS	40038					
37 NO:	INL	30012					
38 NO:	INS	20353			<u> </u>		
39 NO:	INS	20019					
40	INS	20019					

41								
NO: 42	INS	20019						
NO: 43	INL	20052	MAM	10007				
NO:	INL	30012	STO	10021				
NO:	INL	30012						
NO:	INL	30012	ADR	10013				
46 NO:	INS	30028	NRV	10008				
47 NO:	INS	20019						
48 NO:	INS	30028						
19 NO:	INS	30028	BON	1002	LMN	10017		
NO:	INS	30028						
51 NO:	INS	40038	INL	20008		-		
52 NO:	INL	30012	KID	20012				
53 NO:	INL	30012				-	-	
54 NO:	INL	40016	_	-				
55 NO:	INS	50047	CON	10007			-	
56 NO:	INL	30012						
57								
NO: 58	INS	20019	INL	10006	LIV	20038	BLO	1004
NO:	INL	30012						
59				ł				
59 NO: 60	INL	30012	INS	20019				
NO:		30012	INS	20019				
NO: 60 NO: 61 NO:	INL			i				
NO: 60 NO: 61 NO: 62 NO:	INL	30012		i				
NO: 60 NO: 61 NO: 62 NO: 63	INL INL INL	30012		i				
NO: 60 NO: 61 NO: 62 NO: 63	INL INL INL	30012 30012 30012		i				
NO: 60 NO: 61 NO: 62 NO: 63 NO: 64 NO: 65	INL INL INL INL INL	30012 30012 30012 30028		i				
NO: 60 NO: 61 NO: 62 NO: 63 NO: 64 NO: 65	INL INL INL INL INL INL	30012 30012 30012 30028 30012		i				
NO: 60 NO: 61 NO: 62 NO: 63 NO: 64 NO: 65 NO:	INL INL INL INL INL INS INL	30012 30012 30012 30028 30012 30028		i				
NO: 60 NO: 61 NO: 62 NO: 64 NO: 65 NO: 66 NO: 67	INL INL INL INL INL INS INL INS	30012 30012 30012 30028 30012 30028 30012		i				
NO: 60 NO: 61 NO: 62 NO: 63 NO: 64 NO: 65 NO: 66 NO: 67 NO: 68	INL INL INL INL INL INS INL INS	30012 30012 30012 30028 30012 30028 30012		i				
NO: 60 NO: 61 NO: 62 NO: 63 NO: 65 NO: 65 NO: 67 NO: 68 NO: 69	INL INL INL INL INS INL INS INL INS INL INS	30012 30012 30012 30028 30012 30028 30012 30028	PNS	10022				
NO: 60 NO: 61 NO: 62 NO: 63 NO: 65 NO: 65 NO: 67 NO: 67 NO: 69 NO:	INL INL INL INL INS INL INS INL INS INL INS INL INS INS	30012 30012 30012 30028 30012 30028 30028 30028 20019	PNS	10022				
NO: 60 NO: 61 NO: 62 NO: 65 NO: 65 NO: 67 NO: 68 NO: 69 NO: 70 NO: 71 NO:	INL INL INL INL INL INS INL INS INL INS INL INS INS INS	30012 30012 30012 30028 30012 30028 30012 30028 20019 30028	PNS	10022				

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				118			
74		.0067				Τ	
NO: 75	INS	30028					
NO: 76	INS	30028	INL	30012	PIT	10029	
NO: 77	INS	20019	_				
NO: 78	INS	20019					
NO: 79	INS	20019					
NO: 80	INS	20019					
NO: 81	INS	20353					
NO: 82	INS	20353					
NO: 83	INS	20353					
NO: 84	INS	20353					
NO: 85	INS	20162					
NO: 86	INS	20162					
NO: 87	INS	30028					
NO:	INS	30028					
NO: 89	INS	30028					
NO: 90	INS	20162					
NO:. 91	INL	60024	INS	10009	PTH	1004	
NO: 92	INT	203	NOS	10287			
NO: 93	INT	203					
NO: 94	INS	17 - .0161	INL	10004			
NO: 95	INS	17 - .0161	INL	10004			
NO: 96	INS	40038	INL	10004			
NO: 97	INS	30028					
NO: 98	INS	30028					
NO: 99	INS	40038	INL	20008			
NO: 100	INS	40038	INL	20008			

INS Intestine, Small

INT Intestine

5 INL Intestine, Large Connective Tissue CON

MSL Muscles

Nervous Tissue

NRV

SKN Skin

10 NOS Nose UNC Mixed Tissues

EYE Eye

OVR Ovary UTR Uterus FTS Fetus KID Kidney

5

There are 2 values for each organ in the format 9 - 0.9999. The first represent the number of occurrences of the gene in the given organ. The 2nd number represents the percentage of the expression of the gene in the given organ.

Based on sequence alignment with the human genome, the following chromosomal locations were assigned.

	,		
Nucleotide	Chromosomal	ORF location	Amino acid
Name	Location		Name
DEX0234_1	3p26.3c	304-125	DEX0234_101
DEX0234_2	5p15.33e	717-523	DEX0234_102
DEX0234_3	6p25.3b	136-285	DEX0234_103
DEX0234_4	6p25.3b		
DEX0234_5	2p25.3g	123-1	DEX0234_104
DEX0234_6	1p36.33b	620-775	DEX0234_105
DEX0234_7	1p36.33b	160-23	DEX0234_106
DEX0234_8	19p13.3j	471-316	DEX0234_107
DEX0234_9	12p13.33d	1-2787	DEX0234_108
DEX0234_10	17p13.3g	58-1113	DEX0234_109
DEX0234_11	2p25.3g	165-2312	DEX0234_110
DEX0234_12	6p25.3b	131-328	DEX0234_111
DEX0234_13	19p13.3j	66-1	DEX0234_112
DEX0234 14	13p13e	19-84	DEX0234_113
DEX0234 15	11p15.5d	157-32	DEX0234_114
DEX0234_16	11p15.5d	32-160	DEX0234_115
DEX0234 17	5p15.33e		
DEX0234_18	4p16.3d	31-117	DEX0234_116
DEX0234 19	8p23.3b	349-56	DEX0234_117
DEX0234 20	19p13.3j	462-872	DEX0234 118
DEX0234 21	19p13.3j	231-386	DEX0234 119
DEX0234 22	16p13.3f	907-638	DEX0234_120
DEX0234 23	12p13.33d	713-904	DEX0234 121
DEX0234 24	2p25.3g	102-1	DEX0234 122
DEX0234 25	2p25.3g	596-1051	DEX0234_123
DEX0234 26	2p25.3g	465-644	DEX0234 124
DEX0234 27	2p25.3g	865-614	DEX0234 125
DEX0234 28	10p15.3d	288-184	DEX0234 126
DEX0234 29			
DEX0234 30	19p13.3j	1117-3705	DEX0234 127
DEX0234 31	19p13.3j	120-683	DEX0234 128
DEX0234_32	15p13e	48-13	DEX0234_129
DEX0234 33	16p13.3f	1938-1609	DEX0234 130
DEX0234 34	9p24.3b	533-916	DEX0234 131
DEX0234 35	2p25.3g	573-809	DEX0234 132
DEX0234 36	8p23.3b	279-109	DEX0234 133
DEX0234_37	20p13f	351-190	DEX0234_134
DEX0234_38	11p15.5d	186-43	DEX0234_135
DEX0234 39	9p24.3b	1-489	DEX0234_136
DEX0234 40	7p22.3d	546-686	DEX0234_137
DEX0234 41	7p22.3d	321-178	DEX0234 138
DEX0234_42	7p22.3d	40-246	DEX0234_139
DEX0234 43	13p13e	39-1	DEX0234 140

DEX0234_44	3p26.3c	326-3	DEX0234_141
DEX0234_45	3p26.3c	304-188	DEX0234_142
DEX0234_46	10p15.3d	343-119	DEX0234_143
DEX0234_47	5p15.33e	2153-1794	DEX0234_144
DEX0234_48	11p15.5d	68-1	DEX0234_145
DEX0234_49	1p36.33b	39-140	DEX0234_146
DEX0234_50	2p25.3g	810-1202	DEX0234_147
DEX0234_51	12p13.33d	375-220	DEX0234_148
DEX0234_52	7p22.3d		
DEX0234_53	13p13e	733-605	DEX0234_149
DEX0234_54	2p25.3g	529-395	DEX0234_150
DEX0234_55	3p26.3c	204-1	DEX0234_151
DEX0234_56	6p25.3b	275-3	DEX0234_152
DEX0234_57	3p26.3c		
DEX0234_58	16p13.3f	107-577	DEX0234_153
DEX0234_59	15p13e	320-388	DEX0234_154
DEX0234_60	1p36.33b		
DEX0234_61	17p13.3g	288-148	DEX0234_155
DEX0234_62	17p13.3g	698-543	DEX0234_156
DEX0234_63	15p13e	1574-1356	DEX0234_157
DEX0234_64	4p15.3d	16-114	DEX0234_158
DEX0234_65	20p13f	334-275	DEX0234_159
DEX0234_66	19p13.3j		
DEX0234_67	17p13.3g	137-1	DEX0234_160
DEX0234_68	8p23.3b	140-1	DEX0234_161
DEX0234_69	17p13.3g	459-872	DEX0234_162
DEX0234_70	2p25.3g	1596-1390	DEX0234_163
DEX0234_71	11p15.5d	52-2	DEX0234_164
DEX0234_72	3p26.3c	466-2820	DEX0234_165
DEX0234_73	17p13.3g	884-1174	DEX0234_166
DEX0234_74	17p13.3g	940-293	DEX0234_167
DEX0234_75	17p13.3g	257-460	DEX0234_168
DEX0234_76	15p13e	2214-1906	DEX0234_169
DEX0234_77	6p25.3b	461-577	DEX0234_170
DEX0234_78	1p36.33b	157-2	DEX0234_171
DEX0234_79	7p22.3d	134-1	DEX0234_172
DEX0234_80	15p13e		
DEX0234_81	7p22.3d	448-549	DEX0234_173
DEX0234_82	9p24.3b	12-101	DEX0234_174
DEX0234_83	3p26.3c	9-71	DEX0234_175
DEX0234_84	5p15.33e	158-394	DEX0234_176
DEX0234_85	10p15.3d	9-122	DEX0234_177
DEX0234_86	20p13f	293-358	DEX0234_178
DEX0234_87	5p15.33e	246-416	DEX0234_179
DEX0234_88	2p25.3g		
DEX0234_89	7p22.3d		
DEX0234_90	Xp22.33f	228-299	DEX 0234 180
DEX0234_91	2p25.3g	590-312	DEX0234_181
DEX0234_92	16p13.3f	56-139	DEX0234_182
DEX0234 93	16p13.3f	90-590	DEX0234_183
DEX0234_94	17p13.3g	222-464	DEX0234_184
DEX0234_95	17p13.3g	105-584	DEX0234_185
DEX0234_96	12p13.33d	1-2787	DEX0234_186
DEX0234_97	2p25.3g	688-398	DEX0234_187
DEX0234_98	10p15.3d	376-233	DEX0234_188
DEX0234_99	4p16.3d	317-2251	DEX0234_189
DEX0234_100	4p16.3d	252-106	DEX0234_190

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Example 2A: Custom Microarray Experiment-Colon Cancer

Custom oligonucleotide microarrays were provided by Agilent Technologies, Inc. (Palo Alto, CA). The microarrays were fabricated by Agilent using their technology for the *in-situ* synthesis of 60mer oligonucleotides (Hughes, et al. 2001, Nature

Biotechnology 19:342-347). The 60mer microarray probes were designed by Agilent, from gene sequences provided by diaDexus, using Agilent proprietary algorithms.

Whenever possible two different 60mers were designed for each gene of interest.

All microarray experiments were two-color experiments and were preformed using Agilent-recommended protocols and reagents. Briefly, each microarray was hybridized with cRNAs synthesized from polyA+ RNA, isolated from cancer and normal tissues, labeled with fluorescent dyes Cyanine3 and Cyanine5 (NEN Life Science Products, Inc., Boston, MA) using a linear amplification method (Agilent). In each experiment the experimental sample was polyA+ RNA isolated from cancer tissue from a single individual and the reference sample was a pool of polyA+ RNA isolated from normal tissues of the same organ as the cancerous tissue (i.e. normal colon tissue in experiments with colon cancer samples). Hybridizations were carried out at 60°C, overnight using Agilent in-situ hybridization buffer. Following washing, arrays were scanned with a GenePix 4000B Microarray Scanner (Axon Instruments, Inc., Union City, CA). The resulting images were analyzed with GenePix Pro 3.0 Microarray Acquisition and Analysis Software (Axon). Two different chip designs were evaluated with 20 overlapping sets of a total of 38 samples, comparing the expression patterns of colon cancer derived polyA+ RNA to polyA+ RNA isolated from a pool of 7 normal colon tissues were analyzed. For Chip1 all 38 samples (23 Ascending colon carcinomas and 15 Rectosigmoidal carcinomas including: 5 stage 1 cancers, 15 stage 2 cancers, 15 stage 3 25 and 2 stage 4 cancers) were analyzed and for Chip2 a subset of 27 of these samples (14 Ascending colon carcinomas and 13 Rectosigmoidal carcinomas including: 3 stage 1 cancers, 9 stage 2 cancers, 13 stage 3 and 2 stage 4 cancers) were assessed.

Data normalization and expression profiling were done with Expressionist software from GeneData Inc. (Daly City, CA/Basel, Switzerland). Gene expression analysis was performed using only experiments that meet certain quality criteria. The quality criteria that experiments must meet are a combination of evaluations performed by the Expressionist software and evaluations performed manually using raw and normalized data. To evaluate raw data quality, detection limits (the mean signal for a replicated

negative control + 2 Standard Deviations (SD)) for each channel were calculated. The detection limit is a measure of non-specific hybridization. Arrays with poor detection limits were not analyzed and the experiments were repeated. To evaluate normalized data quality, positive control elements included in the array were utilized. These array features should have a mean ratio of 1 (no differential expression). If these features have a mean ratio of greater than 1.5-fold up or down, the experiments were not analyzed further and were repeated. In addition to traditional scatter plots demonstrating the distribution of signal in each experiment, the Expressionist software also has minimum thresholding criteria that employ user defined parameters to identify quality data. Only those features that meet the threshold criteria were included in the filtering and analyses carried out by Expressionist. The thresholding settings employed require a minimum area percentage of 60% [(% pixels > background + 2SD)-(% pixels saturated)], and a minimum signal to noise ratio of 2.0 in both channels. By these criteria, very low expressors and saturated features were not included in analysis.

Relative expression data was collected from Expressionist based on filtering and clustering analyses. Up- and down-regulated genes were identified using criteria for percentage of valid values obtained, and the percentage of experiments in which the gene is up- or down-regulated. These criteria were set independently for each data set, depending on the size and the nature of the data set. The results for the statistically significant upregulated and downregulated genes are shown in Table 1 and Table 2. The first three columns of each table contain information about the sequence itself (Oligo ID, Parent ID, and Patent#), the next 3 columns show the results obtained. "%valid' indicates the percentage of unique experiments total (n=38 for Chip1, n=27 for Chip2) in which a valid expression value was obtained, "%up' indicates the percentage of experiments in which up-regulation of at least 2-fold was observed, and '%down' indicates the percentage of the experiments in which down-regulation of at least 2-fold was observed. The last column in each table describes the location of the microarray probe (oligo) relative to the parent sequence.

Table 1. Colon Microarray expression data.

Patent	ParentID	OligoID	%valid n=38	%down n=38	%valid down n=38	Oligo Start Pos in Par Seq	Oligo Stop Pos in Par Seq
DEX0234_11	11855	26273	94.7	68.4	72.2	158	217
DEX0234_11	11855	26272	92.1	60.5	65.7	209	268
DEX0234_15	11865	23559	13.2	2.6	20	201	260

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Patent	ParentID	OligoID	%valid	%down	%valid	Oligo	Oligo
			n=38	n=38	down	Start	Stop
					n=38	Pos in	Pos in
		l .	1	1		Par Seq	Par Seq
DEX0234_16	11865	23558	86.8	86.8	100	611	670
DEX0234_27,	11897	38407	100	68.4	68.4	1092	1151
DEX0234_24		1					
DEX0234_27	11897	38408	100	63.2	63.2	1051	1110
DEX0234_52	11696	27384	94.7	63.2	66.7	968	1027
DEX0234_52	11696	27383	100	63.2	63.2	1028	1087
DEX0234_60	11848	34228	65.8	55.3	84	1642	1701
DEX0234 60	11848	34229	52.6	28.9	55	1245	1304

Table 2. Colon Microarray Tissue Descriptions.

Experiment			Region			
Name	Renamed	Tissue ID	affected	Stage	Grade	TNM
CNA1C038NMP		BS993038A		1	1	T1N0M0
CNA1C089NMP		S9914089A		1	1 or 2	T1N0M0
CNA1C108NMP		9706C108RA		1	2	T2N0M0
CNA1C281NMP		S992281A		1	1	T2N0M0
CNA1C795NMP		S9915795A		1	1 or 2	T1N0M0
CNA2C012NMP		9608B012		2	2-3	T4N0M0
CNA2C062NMP		S993062C		2	1 or 2	T3N0M0
CNA2C074NMP		9709C074RA		2	2-3	T3N0M0
CNA2C382NMP		4005382A3		2	2	T2N0M0
CNA2C56CNMP		1056C		2	2	T3N0M0
CNA2C677NMP		4005677A1		2	3	T3N0M0
CNA2C695NMP		4004695A9		2	2	T3N0M0
CNA2C821NMP		S9914821A		2	2	T3N0M0
CNA3C003NMP		9610B003		3	3	T3M1M0
CNA3C004NMP		9707C004GB		3	2	T3NIM0
CNA3C032NMP		S9921032A		3	2	T3N1M0
CNA3C068NMP		9706C068RA		3	2 or 3	T3N1M0
CNA3C401NMP		S9819401A		3	2	T2N1M0
CNA3C720NMP		S993720A		3	3	T1N2M0
CNA3C806NMP	CNA4C806NMP	S9915806A		4	2	T3N1M1
CNA3C810NMP		BS986810A		3	2	T41N1M0
CNA4C005NMP		9706C005RA		4	3	T3N2M1
CNA4C006NMP		9609B006		4	2 or 3	T3N2M1
CNR2C020NMP		9408C020R		2	2	T3N0M0
CNR2C024NMP		9704C024RA		2	2	T3N0M0
CNR2C036ANMP		9705C036A		2	2	T3N0M0
CNR2C086NMP		9707C086B		2	2	T3N0M0
CNR2C162NMP	i	9406C162R		2	2	T4N0M0
CNR2C196NMP		S9820196A		2	2	T3N0M0
CNR2C404NMP		S9819404A		2	2	T4N0M0
CNR3C006BNMP		9702B006B		3	2	T3N1M0
CNR3C014NMP		9707C014RA		3	2	T3N2M0
CNR3C022NMP		9611B022F		3	2	T3N1M0
CNR3C036NMP		9706C036RA		3	2	T3N2M0
CNR3C053NMP		9409C053R		3	2	T3N1M0
CNR3C091NMP	CNR4C091NMP	9703C091R		4	3	T3N1M1
CNR3C457NMP		4004457A3		3	3	T3N1M0
CNR3CC98NMP		1194C98		3	2	T3N1M0

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Example 2B : Relative Quantitation of Gene Expression

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Real-Time quantitative PCR with fluorescent Taqman probes is a quantitation detection system utilizing the 5'-3' nuclease activity of Taq DNA polymerase. The method uses an internal fluorescent oligonucleotide probe (Taqman) labeled with a 5' reporter dye and a downstream, 3' quencher dye. During PCR, the 5'-3' nuclease activity of Taq DNA polymerase releases the reporter, whose fluorescence can then be detected by the laser detector of the Model 7700 Sequence Detection System (PE Applied Biosystems, Foster City, CA, USA). Amplification of an endogenous control is used to standardize the amount of sample RNA added to the reaction and normalize for Reverse Transcriptase (RT) efficiency. Either cyclophilin, glyceraldehyde-3-phosphate dehydrogenase (GAPDH), ATPase, or 18S ribosomal RNA (rRNA) is used as this endogenous control. To calculate relative quantitation between all the samples studied, the target RNA levels for one sample were used as the basis for comparative results (calibrator). Quantitation relative to the "calibrator" can be obtained using the standard curve method or the comparative method (User Bulletin #2: ABI PRISM 7700 Sequence Detection System).

The tissue distribution and the level of the target gene are evaluated for every sample in normal and cancer tissues. Total RNA is extracted from normal tissues, cancer tissues, and from cancers and the corresponding matched adjacent tissues. Subsequently, first strand cDNA is prepared with reverse transcriptase and the polymerase chain reaction is done using primers and Taqman probes specific to each target gene. The results are analyzed using the ABI PRISM 7700 Sequence Detector. The absolute numbers are relative levels of expression of the target gene in a particular tissue compared to the calibrator tissue.

One of ordinary skill can design appropriate primers. The relative levels of expression of the CSNA versus normal tissues and other cancer tissues can then be determined. All the values are compared to normal thymus (calibrator). These RNA samples are commercially available pools, originated by pooling samples of a particular tissue from different individuals.

The relative levels of expression of the CSNA in pairs of matching samples and 1 cancer and 1 normal/normal adjacent of tissue may also be determined. All the values are compared to normal thymus (calibrator). A matching pair is formed by mRNA from the cancer sample for a particular tissue and mRNA from the normal adjacent sample for that same tissue from the same individual.

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In the analysis of matching samples, the CSNAs that show a high degree of tissue specificity for the tissue of interest. These results confirm the tissue specificity results obtained with normal pooled samples. Further, the level of mRNA expression in cancer samples and the isogenic normal adjacent tissue from the same individual are compared.

This comparison provides an indication of specificity for the cancer stage (e.g. higher levels of mRNA expression in the cancer sample compared to the normal adjacent).

Altogether, the high level of tissue specificity, plus the mRNA overexpression in matching samples tested are indicative of SEQ ID NO: 1-100 being a diagnostic marker for cancer.

10 Example 3: Protein Expression

The CSNA is amplified by polymerase chain reaction (PCR) and the amplified DNA fragment encoding the CSNA is subcloned in pET-21d for expression in E. coli. In addition to the CSNA coding sequence, codons for two amino acids, Met-Ala, flanking the NH₂-terminus of the coding sequence of CSNA, and six histidines, flanking the 15 COOH-terminus of the coding sequence of CSNA, are incorporated to serve as initiating Met/restriction site and purification tag, respectively.

An over-expressed protein band of the appropriate molecular weight may be observed on a Coomassic blue stained polyacrylamide gel. This protein band is confirmed by Western blot analysis using monoclonal antibody against 6X Histidine tag.

Large-scale purification of CSP is achieved using cell paste generated from 6-liter bacterial cultures, and purified using immobilized metal affinity chromatography (IMAC). Soluble fractions that are separated from total cell lysate were incubated with a nickle chelating resin. The column is packed and washed with five column volumes of wash buffer. CSP is eluted sterwise with various concentration imidazole buffers.

25 Example 4: Fusion Proteins

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The human Fc portion of the IgG molecule can be PCR amplified, using primers that span the 5' and 3' ends of the sequence described below. These primers also should have convenient restriction enzyme sites that will facilitate cloning into an expression vector, preferably a mammalian expression vector. For example, if pC4 (Accession No. 209646) is used, the human Fc portion can be ligated into the BamHI cloning site. Note that the 3' BamHI site should be destroyed. Next, the vector containing the human Fc portion is re-restricted with BamHI, linearizing the vector, and a polynucleotide of the

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present invention, isolated by the PCR protocol described in Example 2, is ligated into this BamHI site. Note that the polynucleotide is cloned without a stop codon, otherwise a fusion protein will not be produced. If the naturally occurring signal sequence is used to produce the secreted protein, pC4 does not need a second signal peptide. Alternatively, if the naturally occurring signal sequence is not used, the vector can be modified to include a heterologous signal sequence. See, e. g., WO 96/34891.

Example 5: Production of an Antibody from a Polypeptide

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In general, such procedures involve immunizing an animal (preferably a mouse) with polypeptide or, more preferably, with a secreted polypeptide-expressing cell. Such 10 cells may be cultured in any suitable tissue culture medium; however, it is preferable to culture cells in Earle's modified Eagle's medium supplemented with 10% fetal bovine serum (inactivated at about 56°C), and supplemented with about 10 g/1 of nonessential amino acids, about 1.000 U/ml of penicillin, and about 100, ug/ml of streptomycin. The splenocytes of such mice are extracted and fused with a suitable myeloma cell line. Any 15 suitable myeloma cell line may be employed in accordance with the present invention: however, it is preferable to employ the parent myeloma cell line (SP20), available from the ATCC. After fusion, the resulting hybridoma cells are selectively maintained in HAT medium, and then cloned by limiting dilution as described by Wands et al., Gastroenterology 80: 225-232 (1981).

The hybridoma cells obtained through such a selection are then assayed to identify clones which secrete antibodies capable of binding the polypeptide. Alternatively, additional antibodies capable of binding to the polypeptide can be produced in a two-step procedure using anti-idiotypic antibodies. Such a method makes use of the fact that antibodies are themselves antigens, and therefore, it is possible to obtain an antibody 25 which binds to a second antibody. In accordance with this method, protein specific antibodies are used to immunize an animal, preferably a mouse. The splenocytes of such an animal are then used to produce hybridoma cells, and the hybridoma cells are screened to identify clones which produce an antibody whose ability to bind to the protein-specific antibody can be blocked by the polypeptide. Such antibodies comprise anti-idiotypic antibodies to the protein specific antibody and can be used to immunize an animal to induce formation of further protein-specific antibodies.

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For the polypeptides of the invention, the following attributes were found, epitopes, post translational modifications, signal peptides and transmembrane domains. Specifically, the Jameson-Wolf methods were used to predicte epitopes were predicted. (Jameson and Wolf, CABIOS, 4(1), 181-186, 1988). Examples of post-translational modifications (PTMs) and other motifs of the CSPs of this invention are listed below. In addition, antibodies that specifically bind such post-translational modifications may be useful as a diagnostic or as therapeutic. The PTMs and other motifs were predicted by using the ProSite Dictionary of Proteins Sites and Patterns (Bairoch et al., Nucleic Acids Res. 25(1):217-221 (1997)), the following motifs, including PTMs, were predicted for the CSPs of the invention. The signal peptides were detected by using the Signal P 2.0, see "Machine learning approaches to the prediction of signal peptides and other protein sorting signals" Henrik Nielsen, Søren Brunak, and Gunnar von Heijne, Protein Engineering 12, 3-9 (1999). Prediction of transmembrane helices in proteins was performed by the application TMHMM 2.0, "currently the best performing transmembrane 15 prediction program", according to authors (A. Krogh, B. Larsson, G. von Heijne, and E. L. L. Sonnhammer, Predicting transmembrane protein topology with a hidden Markov model: Application to complete genomes. Journal of Molecular Biology, 305(3):567-580. January 2001). S. Moller, M.D.R. Croning, R. Apweiler "Evaluation of methods for the prediction of membrane spanning regions" Bioinformatics, 17(7):646-653, July 2001. E. L.L. Sonnhammer, G. von Heijne, and A. Krogh. A hidden Markov model for predicting 20 transmembrane helices in protein sequences in J. Glasgow, T. Littlejohn, F. Major, R. Lathrop, D. Sankoff, and C. Sensen, editors, Proceedings of the Sixth International Conference on Intelligent Systems for Molecular Biology, pages 175-182, Menlo Park, CA, 1998. AAAI Press. Using the PSORT II program, the following cellular localizations 25 and the k nearest neighbors classifier values were determined (Paul Horton and Kenta Nakai, Better Prediction of Protein Cellular Localization Sites with the k Nearest Neighbors Classifier, Intelligent Systems for Molecular Biology 5 147-152 (1997). In the table below are the following: PTM and other motifs (type, amino acid residue locations); Amino acid location and antigenic index (location, AI score, length); TM (number of

30 membrane domain, topology in orientation and position).

	I = =				
name	DEX0234		DEX0234.aa.ptm	DEX0234	
	.aa.ai	234.a		.aa.sp	aa.tmhm
		a.pst			
DEX0234_101	Ĭ	nuc	Ck2_Phospho_Site 27-30;32-35;	28 0.945	
	1		Cytochrome_C 36-41;	0.628	
	1		Pkc_Phospho_Site 27-29;		
DEX0234 102		cyt	Asn_Glycosylation 36-39;		
		· ·	Pkc_Phospho_Site 16-18;		
DEX0234_103		nuc			
DEX0234 104		nuc			
DEX0234_105		pla			
					4 14 5 5 4
DEX0234_106		pla	Myristyl 18-23;31-36;		1 i12-34o
DEX0234_107	i	cyt	Myristyl 14-19;38-43;		
			Pkc_Phospho_Site 20-22;30-32;		
DEX0234_108	53-65	cyt	Amidation 699-702;		
_	1.23 13		Asn Glycosylation 83-86; 120-		
			123:161-164:198-201:239-242:276-		
			279;373-376;902-905; Atp_Gtp_A		
	ĺ		527-534; Camp_Phospho_Site 701-		
			704;876-879; Ck2 Phospho Site		
			11-14:70-73:91-94:169-172:247-		
			250;286-289;376-379;525-528;609-		
	1		612:791-794:809-812:842-845:		
1	1		Myristyl 127-132;205-210;282-		
			287;456-461;471-476;625-630;749-		
			754;786-791;866-871;		
	1		Pkc_Phospho_Site 84-86;130-		
	l		132;162-164;208-210;240-242;414-		
	1		416;540-542;681-683;754-756;770-	1	
	ı		772;791-793;835-837;844-846;848-		
	i		850; Transferrin 1 146-154;224-		
			232;		
	744-755		232,		
	1.19 12	ľ			
	864-880				
1	1.11 17 564-581	1			
	1.06 18				
1	588-604	1			
1	1.06 17	1			
1	435-460	l			
1	1.03 26	1			
1	697-709	1			
1	1.02 13	l			
1	271-281	l			
	1.02 11				
DEX0234_109	335-344	ves	Asn_Glycosylation 228-231;		
	1.06 10		Camp_Phospho_Site 125-128;		
		l	Ck2_Phospho_Site 60-63;		
		l	Leucine_Zipper 205-226; Myristyl		
		l	66-71;112-117;163-168;346-351;		
	1	l	Pkc_Phospho_Site 109-111;124-		
		l	126;169-171;		
	267-277	1		1	
	1.05 11				

			129		
DEX0234_110	85-94 1.14 10 495-536 1.07 42 354-364 1.06 11 382-392 1.05 11 18-36 1.03 19	pla	Amidation 373-376; Asn_Glycosylation 103-106;157- 160;174-177;264-267;452-455;498- 501;539-542;551-554;994-597; Ck2_Phospho_Site 40-43;64-67:85- 88,765-268;287-290;339-342;403- 406;481-484;507-510, Myristyl 144- 149;354-359;383-388;419-424;511- 16;544-549; Pkc_Phospho_Site 54- 56;87-89;108-110;180-182;224- 226;355-357;366-388;474-476;507- 509;515-517;541-543;599-601;608- 107;12-714; Trif_Mgfr_1489-522; Tyr_Phospho_Site 139-147;292- 298; Zinc_Protease 402-411;	17 0.956 0.866	1 0673-695i
DEX0234_111	1.03 19	nuc	Amidation 11-14; Ck2_Phospho_Site 49-52; Pkc_Phospho_Site 11-13;		
DEX0234_112		cyt	Myristyl 12-17;		
DEX0234_113		cyt	Tyr_Phospho_Site 12-19;		
DEX0234_114	8-28 1.00 21	nuc	Asn_Glycosylation 35-38; Ck2_Phospho_Site 10-13;12-15; Myristyl 28-33; Pkc_Phospho_Site 7-9;12-14;		
DEX0234_115		nuc	Myristyl 27-32;		
DEX0234_116		cyt	Ck2_Phospho_Site 19-22;		
DEX0234_117		pla	12; Pkc_Phospho_Site 40-42;		2 i13- 35o64-86i
DEX0234_118	18-35 1.17 18	exc	Camp_Phospho_Site 20-23; Ck2_Phospho_Site 64-67; Myristyl 4-9;37-42; Pkc_Phospho_Site 27- 29;		
DEX0234_119		cyt	Myristyl 43-48;		
DEX0234_120		exc	Ck2_Phospho_Site 62-65; Pkc_Phospho_Site 83-85;	15 0.953 0.880	
DEX0234_121		nuc	Camp_Phospho_Site 45-48; Myristyl 47-52; Pkc_Phospho_Site 55-57;		
DEX0234_122		nuc	Asn_Glycosylation 30-33;		
DEX0234_123	117-137 1.10 21	nuc	Ck2_Phospho_Site 14-17;36-39; Myristyl 46-51;113-118; Pkc_Phospho_Site 11-13;94- 96;100-102;132-134;		
DEX0234_124		pla	Myristyl 20-25; Pkc_Phospho_Site 36-38; Prokar_Lipoprotein 15-25;	23 0.996 0.908	

DEX0234_125	ĺ	exc	Ck2_Phospho_Site 76-79; Myristyl	
		ŀ	40-45; Pkc_Phospho_Site 18-20;76-	
			78;	
DEX0234_126		nuc		
DEX0234_127	445-476	pla	Amidation 682-685;	1 o803-825i
	1.29 32		Asn_Glycosylation 199-202;530-	
			533;617-620;830-833; Atp_Gtp_A	
	1		464-471; Camp_Phospho_Site 613-	
			616; Ck2_Phospho_Site 45-48;80-	
			83;101-104;116-119;150-153;302-	
			305;327-330;336-339;376-379;395-	
		l	398;433-436;632-635;765-768;772-	
		İ	775;824-827;842-845; Glycosaminoglycan 747-750;	
			Myristyl 217-222;238-243;390-	
			395;431-436;464-469;585-590;	
			Pkc_Phospho_Site 185-187;213-	
1			215;225-227;302-304;409-411;445-	
			447;473-475;501-503;611-613;673-	
			675;682-684;857-859;	
			Tyr_Phospho_Site 684-691;685-	
			691; Zinc_Finger_C2h2 437-	
			457;465-485;493-513;521-541;549-	
1		l	569;577-597;605-625;633-653;661-	
			681;	
	765-800		·	
	1.28 36			
	251-263			
	1.27 13			
	76-86	ŀ		
	1.24 11			
	482-508			
	1.17 27			
	648-666 1.16 19			
1	510-582			
	1.13 73			
	584-642			
	1.12 59			
1	39-74			
	1.03 36			
DEX0234_128	76-86	nuc	Ck2_Phospho_Site 45-48;80-	
_	1.24 11		83;101-104;116-119;150-153;	
		l	Pkc_Phospho_Site 185-187;	
	39-74	l		
	1.03 36			
DEX0234_129			Tyr_Phospho_Site 3-11;	
DEX0234_130	4-22 1.10	nuc	Asn_Glycosylation 4-7;	
	19		Ck2_Phospho_Site 13-16;34-37;	
		1	Myristyl 29-34;84-89;	
			Pkc_Phospho_Site 70-72;79-81;	
DEX0234_131		ves	Myristyl 44-49;57-62;61-66;90-95;	2 074-
				96i103-
				1250
DEX0234_132	33-50	nuc	Camp_Phospho_Site 58-61;	
	1.01 18		Pkc_Phospho_Site 56-58;	

DEX0234_133		mit	Myristyl 27-32;36-41;40-45; Pkc_Phospho_Site 48-50;		1 o22-44i
DEX0234_134		cyt	Myristyl 9-14;13-18; Pkc Phospho Site 14-16;		
DEX0234 135		cyt	Myristyl 21-26:		
DEX0234 136	51-76	ves			
	1.06 26		79;135-140; Pkc_Phospho_Site 8- 10;52-54;154-156;		
DEX0234_137	19-35 1.16 17	exc	Asn_Glycosylation 43-46; Myristyl 26-31;	15 0.992 0.951	
DEX0234_138	18-36 1.08 19	cyt	Asn_Glycosylation 29-32;		
DEX0234_139	21-51	nuc	Ck2_Phospho_Site 28-31;42-45;		
	1.16 31		Myristyl 13-18;38-43;50-55; Pkc_Phospho_Site 35-37;41-43;42- 44:		
DEX0234 141	49-68	nuc	Asn Glycosylation 65-68:		
52.0251_111	1.17 20	""	Ck2_Phospho_Site 56-59;		
			Pkc_Phospho_Site 25-27;30-32;56- 58;84-86;		
DEX0234_142		cyt	Pkc_Phospho_Site 7-9;		
DEX0234_143	41-61	cyt			
	1.16 21		42-47;46-51;55-60;59-64;63-68;64- 69;		
DEX0234_144		pla	Ck2_Phospho_Site 45-48; Myristyl 5-10; Pkc_Phospho_Site 54-56;82- 84;91-93;	52 0.996 0.764	1 13-350
DEX0234_145		nuc	Ck2_Phospho_Site 13-16; Myristyl 4-9; Pkc_Phospho_Site 8-10;20-22;		
DEX0234_146		cyt	Ck2_Phospho_Site 8-11; Pkc_Phospho_Site 28-30;		
DEX0234_147	96-106 1.34 11	nuc	Ck2_Phospho_Site 77-80; Myristyl 26-31; Pkc_Phospho_Site 11-13;		
DEX0234_148	16-26 1.04 11	nuc	Ck2_Phospho_Site 2-5; Pkc_Phospho_Site 2-4;34-36;		
DEX0234_149	26-40 1.16 15	nuc			
DEX0234_150		cyt	Pkc_Phospho_Site 2-4;		
DEX0234_151	34-58 1.14 25	exc	Ck2_Phospho_Site 42-45;50-53; Pkc_Phospho_Site 42-44;	38 0.962 0.620	
DEX0234_152	13-24	nuc	Asn_Glycosylation 22-25;		
	1.19 12		Ck2_Phospho_Site 10-13;33-36;		
			Pkc_Phospho_Site 6-8; Tyr_Phospho_Site 76-83;		
DEX0234_153	96-136	mit	Amidation 111-114;	24 0.924	
	1.03 41		Ck2_Phospho_Site 41-44;	0.693	
			Glycosaminoglycan 56-59;60-63; Myristyl 17-22;19-24;21-26;36-		
			41;39-44;57-62;59-64;61-66;91-		
			96;117-122;118-123;122-127;130-		
ł			135;145-150; Pkc_Phospho_Site 25-		
BE100004 4			27; Prokar_Lipoprotein 14-24;		
DEX0234_154		cyt	Ck2_Phospho_Site 17-20;		

DEX0234_155		cyt	Asn_Glycosylation 13-16;		
			Ck2_Phospho_Site 15-18;		
DEX0234_156		cyt	Pkc_Phospho_Site 35-37;		
DEX0234_157		nuc	Asn_Glycosylation 25-28;		
_			Ck2_Phospho_Site 66-69;		
		l	Pkc_Phospho_Site 11-13;45-47;		
			Prokar_Lipoprotein 49-59;		
DEX0234_158		cyt	-		
DEX0234 160	9-18 1.05	nuc	Ck2_Phospho_Site 37-40;		
	10		Prokar_Lipoprotein 17-27;		
DEX0234 161		nuc	Ck2 Phospho Site 8-11;		
			Pkc_Phospho_Site 21-23;		
DEX0234_162		nuc	Amidation 15-18; Asn_Glycosylation		
_			73-76; Camp_Phospho_Site 61-64;		
			Myristyl 30-35;92-97;123-128;		İ
			Pkc_Phospho_Site 64-66;		
DEX0234 163		cyt	Amidation 62-65; Myristyl 13-18;		
DEX0234 164		-70	Asn_Glycosylation 2-5;		
			Pkc_Phospho_Site 5-7;		l
DEX0234_165	241-251	nuc	Asn_Glycosylation 465-468;		
_	1.20 11		Camp_Phospho_Site 227-230;		
			Ck2_Phospho_Site 3-6;31-34;38-		
			41;132-135;137-140;239-242;484-		
1			487;635-638;691-694;726-729;		
			Myristyl 46-51;97-102;167-172;183-	1	
1			188;225-230;229-234;449-454;		
i			Pkc_Phospho_Site 141-143;171-		1
			173;226-228;333-335;407-409;429-		
			431;501-503;508-510;654-656;		
	164-181		Zinc_Finger_C2h2 16-37;		
	1.17 18				
	341-354				
	1.16 14				
	202-231				
	1.13 30				
	720-735				
	1.10 16				
1	679-702				
	1.09 24				l
	447-459				l
	1.02 13				
DEX0234_166	58-75	cyt	Ck2_Phospho_Site 21-24; Myristyl		
	1.22 18		58-63; Pkc_Phospho_Site 26-28;		
DEX0234_167	115-131	nuc	Amidation 206-209;		1 o92-114i
	1.14 17		Ck2_Phospho_Site 26-29; Myristyl		
			16-21;20-25;47-52;105-110;130-		l
			135;155-160;158-163;		l
			Pkc_Phospho_Site 17-19;64- 66;114-116;124-126;125-127;162-		l
			164;		l
1	54-80		104,		
	1.09 27		_		

		133			
		Myristyl 8-13; Pkc_Phospho_Site 2- 4;13-15;54-56;	nuc		DEX0234_168
		Amidation 90-93; Myristyl 40-45;75- 80;78-83;	nuc		DEX0234_169
		Amidation 19-22; Asn_Glycosylation 17-20; Ck2_Phospho_Site 6-9; Pkc_Phospho_Site 19-21;	nuc	11-28 1.08 18	DEX0234_170
		Asn_Glycosylation 22-25;	nuc		DEX0234_171
1 (13-350	25 0.986 0.919	Myristyl 21-26; Pkc_Phospho_Site 9-11;31-33;	exc		DEX0234_172
		Pkc_Phospho_Site 28-30;	cyt		DEX0234_173
		Pkc_Phospho_Site 17-19;	cyt	13-25 1.18 13	DEX0234_174
			exc		DEX0234_175
		Camp_Phospho_Site 30-33;	nuc		DEX0234_176
		Pkc_Phospho_Site 2-4;	cyt	22-34 1.03 13	DEX0234_177
			nuc		DEX0234_178
		Myristyl 34-39; Pkc_Phospho_Site 30-32;	cyt		DEX0234_179
		Pkc_Phospho_Site 19-21;	cyt		DEX0234_180
		Ck2_Phospho_Site 33-36; Myristyl 46-51; Pkc_Phospho_Site 49-51;63- 65;89-91;	nuc		DEX0234_181
			nuc		DEX0234_182
1 o31-53		Ck2_Phospho_Site 87-90;141-144; Myristyl 115-120; Pkc_Phospho_Site 9-11;66-68;87-89;104-106; Tyr_Phospho_Site 89-96;	pla	104-113 1.01 10	DEX0234_183
	21 0.945 0.793	Asn_Glycosylation 62-65; Ck2_Phospho_Site 47-50; Myristyl 39-44;43-48;57-62;	exc	29-70 1.04 42	DEX0234_184
		Asn_Glycosylation 62-65;106-109; Camp_Phospho_Site 55-58;104- 107; Ck2_Phospho_Site 41-44;50- 53; Pkc_Phospho_Site 25-27;41- 43;54-56;118-120;	nuc	103-112 1.36 10 14-30 1.20 17	DEX0234_185
		Amidation 699-702; Asn. Glycosylation 33-86;120- 123;161-164;198-201;239-242;276- 279;373-376;902-905; Atp. Gtp. A 527-534; Camp_Phospho_Site 701- 704;876-879; Ck2_Phospho_Site 11-14;70-73;91-94;169-172;247- 250;286-289;376-379;255-525;60- 612;791-794;809-81;2942-845; Myristyl 127-132;205-210;282- 287;456-641;741-7476;625-630;749- 754;786-791;866-871; Pkc_Phospho_Site 84-86;130- 132;162-164;208-210;240-242;414-	cyt	53-65 1.23 13	DEX0234_186

			772;791-793;835-837;844-846;848-		
			850; Transferrin_1 146-154;224-		
			232;		
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	744 755				
	744-755				l
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	864-880				
	1.11 17				
	564-581				
	1.06 18				
	588-604				1
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	435-460			i	
	1.03 26				
	697-709			1	1
1	1.02 13			I	
	271-281			l	ŀ
	1.02 11				
DEX0234_187	40-62	nuc	Myristyl 21-26; Pkc_Phospho_Site		
	1.14 23		82-84;		
DEX0234 188		pla	Amidation 44-47; Pkc Phospho Site		1 i25-43o
			9-11:		- 1.00
DEX0234_189	526-538	pla	Amidation 99-102;129-132;		1 i139-161o
	1.29 13		Asn_Glycosylation 46-49;510-		
			513;520-523;556-559;		
			Ck2_Phospho_Site 83-86;122-		
1			125;214-217;242-245;263-266;333-		
			336;357-360;376-379;380-383;392-		!
			395;439-442;455-458;459-462;465-		
			468;487-490;498-501;512-515;516-		
			519;543-546;552-555;558-561;604-		
			607;611-614;628-631; Myristyl 50-		
			55;57-62;90-95;179-184;260-		
			265;264-269;267-272;324-329;372-		
			377;472-477;548-553;	l	
1			Pkc_Phospho_Site 36-38;119-	I	1
1			121;122-124;242-244;332-334;484-	I	1
1 1			486;560-562;604-606;	I	1
	235-268			l	l
	1.27 34			l	ľ
1	416-431			I	l
	1.24 16			l	l
				I	l
1	445-473			l	l
	1.14 29			I	l
1	329-341			I	l
	1.10 13			l	l
	545-584			l	l
1	1.08 40			I	l
				l	l
	83-95			l	l
1					
	1.08 13				
	1.08 13 163-210				
	163-210				

135

	1.05 17			
DEX0234_190		cyt		

Example 6: Method of Determining Alterations in a Gene Corresponding to a Polynucleotide

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RNA is isolated from individual patients or from a family of individuals that have a phenotype of interest. cDNA is then generated from these RNA samples using protocols known in the art. See, Sambrook (2001), supra. The cDNA is then used as a template for PCR, employing primers surrounding regions of interest in SEQ ID NO: 1-100. Suggested PCR conditions consist of 35 cycles at 95°C for 30 seconds; 60-120 seconds at 52-58°C; and 60-120 seconds at 70°C, using buffer solutions described in Sidransky et al., Science 252(5006): 706-9 (1991). See also Sidransky et al., Science 278(5340): 1054-9 (1997).

PCR products are then sequenced using primers labeled at their 5° end with T4 polynucleotide kinase, employing SequiTherm Polymerase. (Epicentre Technologies). The intron-exon borders of selected exons is also determined and genomic PCR products analyzed to confirm the results. PCR products harboring suspected mutations are then cloned and sequenced to validate the results of the direct sequencing. PCR products is cloned into T-tailed vectors as described in Holton et al., Nucleic Acids Res., 19: 1156 (1991) and sequenced with T7 polymerase (United States Biochemical). Affected individuals are identified by mutations not present in unaffected individuals.

Genomic rearrangements may also be determined. Genomic clones are nick-translated with digoxigenin deoxyuridine 5' triphosphate (Boehringer Manheim), and FISH is performed as described in Johnson et al., Methods Cell Biol. 35: 73-99 (1991). Hybridization with the labeled probe is carried out using a vast excess of human cot-1 DNA for specific hybridization to the corresponding genomic locus.

Chromosomes are counterstained with 4,6-diamino-2-phenylidole and propidium iodide, producing a combination of C-and R-bands. Aligned images for precise mapping are obtained using a triple-band filter set (Chroma Technology, Brattleboro, VT) in combination with a cooled charge-coupled device camera (Photometrics, Tucson, AZ) and variable excitation wavelength filters. Id. Image collection, analysis and chromosomal fractional length measurements are performed using the ISee Graphical Program System. (Inovision Corporation, Durham, NC.) Chromosome alterations of the genomic region

hybridized by the probe are identified as insertions, deletions, and translocations. These alterations are used as a diagnostic marker for an associated disease.

Example 7: Method of Detecting Abnormal Levels of a Polypeptide in a Biological Sample

Antibody-sandwich ELISAs are used to detect polypeptides in a sample, preferably a biological sample. Wells of a microtiter plate are coated with specific antibodies, at a final concentration of 0.2 to 10 ug/ml. The antibodies are either monoclonal or polyclonal and are produced by the method described above. The wells are blocked so that non-specific binding of the polypeptide to the well is reduced. The coated wells are then incubated for > 2 hours at RT with a sample containing the polypeptide. Preferably, serial dilutions of the sample should be used to validate results. The plates are then washed three times with deionized or distilled water to remove unbound polypeptide. Next, 50 µl of specific antibody-alkaline phosphatase conjugate, at a concentration of 25-400 ng, is added and incubated for 2 hours at room temperature. The plates are again washed three times with deionized or distilled water to remove unbound conjugate. 75 µl of 4-methylumbelliferyl phosphate (MUP) or p-nitrophenyl phosphate (NPP) substrate solution are added to each well and incubated 1 hour at room temperature.

The reaction is measured by a microtiter plate reader. A standard curve is prepared, using serial dilutions of a control sample, and polypeptide concentrations are plotted on the X-axis (log scale) and fluorescence or absorbance on the Y-axis (linear scale). The concentration of the polypeptide in the sample is calculated using the standard curve.

Example 8: Formulating a Polypentide

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The secreted polypeptide composition will be formulated and dosed in a fashion consistent with good medical practice, taking into account the clinical condition of the individual patient (especially the side effects of treatment with the secreted polypeptide alone), the site of delivery, the method of administration, the scheduling of administration, and other factors known to practitioners. The "effective amount" for purposes herein is thus determined by such considerations.

As a general proposition, the total pharmaceutically effective amount of secreted polypeptide administered parenterally per dose will be in the range of about 1, µg/kg/day to 10 mg/kg/day of patient body weight, although, as noted above, this will be subject to therapeutic discretion. More preferably, this dose is at least 0.01 mg/kg/day, and most

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preferably for humans between about 0.01 and 1 mg/kg/day for the hormone. If given continuously, the secreted polypeptide is typically administered at a dose rate of about 1 µg/kg/hour to about 50 mg/kg/hour, either by 1-4 injections per day or by continuous subcutaneous infusions, for example, using a mini-pump. An intravenous bag solution may also be employed. The length of treatment needed to observe changes and the interval following treatment for responses to occur appears to vary depending on the desired effect.

Pharmaceutical compositions containing the secreted protein of the invention are administered orally, rectally, parenterally, intracistemally, intravaginally, intraperitoneally, topically (as by powders, ointments, gels, drops or transdermal patch), bucally, or as an oral or nasal spray. "Pharmaceutically acceptable carrier" refers to a non-toxic solid, semisolid or liquid filler, diluent, encapsulating material or formulation auxiliary of any type. The term "parenteral" as used herein refers to modes of administration which include intravenous, intramuscular, intraperitoneal, intrasternal, subcutaneous and infusion.

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The secreted polypeptide is also suitably administered by sustained-release systems. Suitable examples of sustained-release compositions include semipermeable polymer matrices in the form of shaped articles, e. g., films, or microcapsules. Sustainedrelease matrices include polylactides (U. S. Pat. No.3,773,919, EP 58,481), copolymers of L-glutamic acid and gamma-ethyl-L-glutamate (Sidman, U. et al., Biopolymers 22: 547-556 (1983)), poly (2-hydroxyethyl methacrylate) (R. Langer et al., J. Biomed, Mater. Res. 15: 167-277 (1981), and R. Langer, Chem. Tech. 12: 98-105 (1982)), ethylene vinyl acetate (R. Langer et al.) or poly-D- (-)-3-hydroxybutyric acid (EP 133,988). Sustainedrelease compositions also include liposomally entrapped polypeptides. Liposomes containing the secreted polypeptide are prepared by methods known per se; DE Epstein et al., Proc. Natl. Acad. Sci. USA 82: 3688-3692 (1985): Hwang et al., Proc. Natl. Acad. Sci. USA 77: 4030-4034 (1980); EP 52,322; EP 36,676; EP 88,046; EP 143,949; EP 142,641; Japanese Pat. Appl. 83-118008; U. S. Pat. Nos. 4,485,045 and 4,544,545; and EP 102,324. Ordinarily, the liposomes are of the small (about 200-800 Angstroms) unilamellar type in which the lipid content is greater than about 30 mol. percent cholesterol, the selected proportion being adjusted for the optimal secreted polypeptide therapy.

For parenteral administration, in one embodiment, the secreted polypeptide is formulated generally by mixing it at the desired degree of purity, in a unit dosage injectable form (solution, suspension, or emulsion), with a pharmaceutically acceptable

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carrier, i. e., one that is non-toxic to recipients at the dosages and concentrations employed and is compatible with other ingredients of the formulation.

For example, the formulation preferably does not include oxidizing agents and other compounds that are known to be deleterious to polypeptides. Generally, the

5 formulations are prepared by contacting the polypeptide uniformly and intimately with liquid carriers or finely divided solid carriers or both. Then, if necessary, the product is shaped into the desired formulation. Preferably the carrier is a parenteral carrier, more preferably a solution that is isotonic with the blood of the recipient. Examples of such carrier vehicles include water, saline, Ringer's solution, and dextrose solution. Non
10 aqueous vehicles such as fixed oils and ethyl oleate are also useful herein, as well as linosomes.

The carrier suitably contains minor amounts of additives such as substances that enhance isotonicity and chemical stability. Such materials are non-toxic to recipients at the dosages and concentrations employed, and include buffers such as phosphate, citrate, succinate, acetic acid, and other organic acids or their salts; antioxidants such as ascorbic acid; low molecular weight (less than about ten residues) polypeptides, e. g., polyarginine or tripeptides; proteins, such as serum albumin, gelatin, or immunoglobulins; hydrophilic polymers such as polyvinylpyrrolidone; amino acids, such as glycine, glutamic acid, aspartic acid, or arginine; monosaccharides, disaccharides, and other carbohydrates including cellulose or its derivatives, glucose, manose, or dextrins; chelating agents such as EDTA; sugar alcohols such as mannitol or sorbitol; counterions such as sodium; and/or nonionic surfactants such as nolvsorbates, poloxamers, or PEG.

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The secreted polypeptide is typically formulated in such vehicles at a concentration of about 0.1 mg/ml to 100 mg/ml, preferably 1-10 mg/ml, at a pH of about 3 to 8. It will be understood that the use of certain of the foregoing excipients, carriers, or stabilizers will result in the formation of polypeptide salts.

Any polypeptide to be used for therapeutic administration can be sterile. Sterility is readily accomplished by filtration through sterile filtration membranes (e. g., 0.2 micron membranes). Therapeutic polypeptide compositions generally are placed into a container having a sterile access port, for example, an intravenous solution bag or vial having a stopper pierceable by a hypodermic injection needle.

Polypeptides ordinarily will be stored in unit or multi-dose containers, for example, sealed ampules or vials, as an aqueous solution or as a lyophilized formulation

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for reconstitution. As an example of a lyophilized formulation, 10-ml vials are filled with 5 ml of sterile-filtered 1 % (w/v) aqueous polypeptide solution, and the resulting mixture is lyophilized. The infusion solution is prepared by reconstituting the lyophilized polypeptide using bacteriostatic Water-for-Injection.

The invention also provides a pharmaceutical pack or kit comprising one or more containers filled with one or more of the ingredients of the pharmaceutical compositions of the invention. Associated with such container (s) can be a notice in the form prescribed by a governmental agency regulating the manufacture, use or sale of pharmaceuticals or biological products, which notice reflects approval by the agency of manufacture, use or sale for human administration. In addition, the polypeptides of the present invention may be employed in confunction with other therapeutic compounds.

Example 9: Method of Treating Decreased Levels of the Polypeptide

It will be appreciated that conditions caused by a decrease in the standard or normal expression level of a secreted protein in an individual can be treated by administering the polypeptide of the present invention, preferably in the secreted form. Thus, the invention also provides a method of treatment of an individual in need of an increased level of the polypeptide comprising administering to such an individual a pharmaceutical composition comprising an amount of the polypeptide to increase the activity level of the polypeptide in such an individual.

For example, a patient with decreased levels of a polypeptide receives a daily dose 0.1-100 ug/kg of the polypeptide for six consecutive days. Preferably, the polypeptide is in the secreted form. The exact details of the dosing scheme, based on administration and formulation, are provided above.

Example 10: Method of Treating Increased Levels of the Polypeptide

Antisense technology is used to inhibit production of a polypeptide of the present invention. This technology is one example of a method of decreasing levels of a polypeptide, preferably a secreted form, due to a variety of etiologies, such as cancer.

For example, a patient diagnosed with abnormally increased levels of a polypeptide is administered intravenously antisense polynucleotides at 0.5, 1.0, 1.5, 2.0 and 3.0 mg/kg day for 21 days. This treatment is repeated after a 7-day rest period if the

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treatment was well tolerated. The formulation of the antisense polynucleotide is provided above.

Example 11: Method of Treatment Using Gene Therapy

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One method of gene therapy transplants fibroblasts, which are capable of expressing a polypeptide, onto a patient. Generally, fibroblasts are obtained from a subject by skin biopsy. The resulting tissue is placed in tissue-culture medium and separated into small pieces. Small chunks of the tissue are placed on a wet surface of a tissue culture flask, approximately ten pieces are placed in each flask. The flask is turned upside down, closed tight and left at room temperature over night. After 24 hours at room temperature, the flask is inverted and the chunks of tissue remain fixed to the bottom of the flask and fresh media (e. g., Ham's F12 media, with 10% FBS, penicillin and streptomycin) is added. The flasks are then incubated at 37°C for approximately one week.

At this time, fresh media is added and subsequently changed every several days. After an additional two weeks in culture, a monolayer of fibroblasts emerge. The monolayer is trypsinized and scaled into larger flasks. pMV-7 (Kirschmeier, P. T. et al., DNA, 7: 219-25 (1988)), flanked by the long terminal repeats of the Moloney murine sarcoma virus, is digested with EcoRI and HindIII and subsequently treated with calf intestinal phosphatase. The linear vector is fractionated on agarose gel and purified, using class heads

The cDNA encoding a polypeptide of the present invention can be amplified using PCR primers which correspond to the 5'and 3'end sequences respectively as set forth in Example 1. Preferably, the 5'primer contains an EcoRI site and the 3'primer includes a HindIII site. Equal quantities of the Moloney murine sarcoma virus linear backbone and the amplified EcoRI and HindIII fragment are added together, in the presence of T4 DNA ligase. The resulting mixture is maintained under conditions appropriate for ligation of the two fragments. The ligation mixture is then used to transform bacteria HB 101, which are then plated onto agar containing kanamycin for the purpose of confirming that the vector has the gene of interest properly inserted.

The amphotropic pA317 or GP+aml2 packaging cells are grown in tissue culture to confluent density in Dulbecco's Modified Eagles Medium (DMEM) with 10% calf serum (CS), penicillin and streptomycin. The MSV vector containing the gene is then added to the media and the packaging cells transduced with the vector. The packaging cells now

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produce infectious viral particles containing the gene (the packaging cells are now referred to as producer cells).

Fresh media is added to the transduced producer cells, and subsequently, the media is harvested from a 10 cm plate of confluent producer cells. The spent media, containing the infectious viral particles, is filtered through a millipore filter to remove detached producer cells and this media is then used to infect fibroblast cells. Media is removed from a sub-confluent plate of fibroblasts and quickly replaced with the media from the producer cells. This media is removed and replaced with fresh media.

If the titer of virus is high, then virtually all fibroblasts will be infected and no selection is required. If the titer is very low, then it is necessary to use a retroviral vector that has a selectable marker, such as neo or his. Once the fibroblasts have been efficiently infected, the fibroblasts are analyzed to determine whether protein is produced.

The engineered fibroblasts are then transplanted onto the host, either alone or after having been grown to confluence on cytodex 3 microcarrier beads.

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Example 12: Method of Treatment Using Gene Therapy-In Vivo

Another aspect of the present invention is using in vivo gene therapy methods to treat disorders, diseases and conditions. The gene therapy method relates to the introduction of naked nucleic acid (DNA, RNA, and antisense DNA or RNA) sequences into an animal to increase or decrease the expression of the polypeptide.

The polynucleotide of the present invention may be operatively linked to a promoter or any other genetic elements necessary for the expression of the polypeptide by the target tissue. Such gene therapy and delivery techniques and methods are known in the art, see, for example, W0 90/11092, W0 98/11779; U. S. Patent No. 5,693,622; 5,705,151; 5,580,859; Tabata H. et al. (1997) Cardiovasc. Res. 35 (3): 470-479, Chao J et al. (1997) Pharmacol. Res. 35 (6): 517-522, Wolff J. A. (1997) Neuromuscul. Disord. 7 (5): 314-318, Schwartz B. et al. (1996) Gene Ther. 3 (5): 405-411, and Tsurumi Y. et al. (1996) Circulation 94 (12): 3281-3290.

The polynucleotide constructs may be delivered by any method that delivers injectable materials to the cells of an animal, such as, injection into the interstitial space of tissues (heart, muscle, skin, colon, liver, intestine and the like). The polynucleotide constructs can be delivered in a pharmaceutically acceptable liquid or aqueous carrier.

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The term "naked" polynucleotide, DNA or RNA, refers to sequences that are free from any delivery vehicle that acts to assist, promote, or facilitate entry into the cell, including viral sequences, viral particles, liposome formulations, lipofectin or precipitating agents and the like. However, the polynucleotides of the present invention may also be delivered in liposome formulations (such as those taught in Felgner P. L. et al. (1995) Ann. NY Acad. Sci. 772: 126-139 and Abdallah B. et al. (1995) Biol. Cell 85 (1): 1-7) which can be prepared by methods well known to those skilled in the art.

The polynucleotide vector constructs used in the gene therapy method are preferably constructs that will not integrate into the host genome nor will they contain sequences that allow for replication. Any strong promoter known to those skilled in the art can be used for driving the expression of DNA. Unlike other gene therapies techniques, one major advantage of introducing naked nucleic acid sequences into target cells is the transitory nature of the polynucleotide synthesis in the cells. Studies have shown that non-replicating DNA sequences can be introduced into cells to provide production of the desired polypeptide for periods of up to six months.

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The polynucleotide construct can be delivered to the interstitial space of tissues within the an animal, including of muscle, skin, brain, colon, liver, spleen, bone marrow. thymus, heart, lymph, blood, bone, cartilage, pancreas, kidney, gall bladder, stomach, intestine, testis, ovary, uterus, rectum, nervous system, eye, gland, and connective tissue. Interstitial space of the tissues comprises the intercellular fluid, mucopolysaccharide matrix among the reticular fibers of organ tissues, elastic fibers in the walls of vessels or chambers, collagen fibers of fibrous tissues, or that same matrix within connective tissue ensheathing muscle cells or in the lacunae of bone. It is similarly the space occupied by the plasma of the circulation and the lymph fluid of the lymphatic channels. Delivery to the interstitial space of muscle tissue is preferred for the reasons discussed below. They may be conveniently delivered by injection into the tissues comprising these cells. They are preferably delivered to and expressed in persistent, non-dividing cells which are differentiated, although delivery and expression may be achieved in non-differentiated or less completely differentiated cells, such as, for example, stem cells of blood or skin fibroblasts. In vivo muscle cells are particularly competent in their ability to take up and express polynucleotides.

For the naked polynucleotide injection, an effective dosage amount of DNA or RNA will be in the range of from about 0.05 µg/kg body weight to about 50 mg/kg body

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weight. Preferably the dosage will be from about 0.005 mg/kg to about 20 mg/kg and more preferably from about 0.05 mg/kg to about 5 mg/kg. Of course, as the artisan of ordinary skill will appreciate, this dosage will vary according to the tissue site of injection. The appropriate and effective dosage of nucleic acid sequence can readily be determined by those of ordinary skill in the art and may depend on the condition being treated and the route of administration. The preferred route of administration is by the parenteral route of injection into the interstitial space of tissues. However, other parenteral routes may also be used, such as, inhalation of an aerosol formulation particularly for delivery to colons or bronchial tissues, throat or mucous membranes of the nose. In addition, naked polynucleotide constructs can be delivered to arteries during angioplasty by the catheter used in the procedure.

The dose response effects of injected polynucleotide in muscle in vivo is determined as follows. Suitable template DNA for production of mRNA coding for polypeptide of the present invention is prepared in accordance with a standard recombinant DNA methodology. The template DNA, which may be either circular or linear, is either used as naked DNA or complexed with liposomes. The quadriceps muscles of mice are then injected with various amounts of the template DNA.

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Five to six week old female and male Balb/C mice are anesthetized by intraperitoneal injection with 0.3 ml of 2.5% Avertin. A 1.5 cm incision is made on the anterior thigh, and the quadriceps muscle is directly visualized. The template DNA is injected in 0.1 ml of carrier in a 1 cc syringe through a 27 gauge needle over one minute, approximately 0.5 cm from the distal insertion site of the muscle into the knee and about 0.2 cm deep. A suture is placed over the injection site for future localization, and the skin is closed with stainless steel clins.

After an appropriate incubation time (e. g., 7 days) muscle extracts are prepared by excising the entire quadriceps. Every fifth 15 um cross-section of the individual quadriceps muscles is histochemically stained for protein expression. A time course for protein expression may be done in a similar fashion except that quadriceps from different mice are harvested at different times. Persistence of DNA in muscle following injection may be determined by Southern blot analysis after preparing total cellular DNA and HIRT supernatants from injected and control mice.

The results of the above experimentation in mice can be use to extrapolate proper dosages and other treatment parameters in humans and other animals using naked DNA.

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Example 13: Transgenic Animals

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The polypeptides of the invention can also be expressed in transgenic animals. Animals of any species, including, but not limited to, mice, rats, rabbits, hamsters, guinea pigs, pigs, micro-pigs, goats, sheep, cows and non-human primates, e. g., baboons, 5 monkeys, and chimpanzees may be used to generate transgenic animals. In a specific embodiment, techniques described herein or otherwise known in the art, are used to

express polypeptides of the invention in humans, as part of a gene therapy protocol.

Any technique known in the art may be used to introduce the transgene (I. e., polynucleotides of the invention) into animals to produce the founder lines of transgenic animals. Such techniques include, but are not limited to, pronuclear microiniection (Paterson et al., Appl. Microbiol. Biotechnol. 40: 691-698 (1994); Carver et al., Biotechnology (NY) 11: 1263-1270 (1993); Wright et al., Biotechnology (NY) 9: 830-834 (1991); and Hoppe et al., U. S. Pat. No. 4,873,191 (1989)); retrovirus mediated gene transfer into germ lines (Van der Putten et al., Proc. Natl. Acad. Sci., USA 82: 6148-6152 15 (1985)), blastocysts or embryos; gene targeting in embryonic stem cells (Thompson et al., Cell 56: 313-321 (1989)); electroporation of cells or embryos (Lo. 1983, Mol Cell, Biol. 3: 1803-1814 (1983)); introduction of the polynucleotides of the invention using a gene gun (see, e. g., Ulmer et al., Science 259: 1745 (1993); introducing nucleic acid constructs into embryonic pleuripotent stem cells and transferring the stem cells back into the blastocyst: and sperm mediated gene transfer (Lavitrano et al., Cell 57: 717-723 (1989), For a review of such techniques, see Gordon, "Transgenic Animals," Intl. Rev. Cytol. 115: 171-229 (1989).

Any technique known in the art may be used to produce transgenic clones containing polynucleotides of the invention, for example, nuclear transfer into enucleated oocytes of nuclei from cultured embryonic, fetal, or adult cells induced to quiescence (Campell et al., Nature 380: 64-66 (1996); Wilmut et al., Nature 385: 810813 (1997)).

The present invention provides for transgenic animals that carry the transgene in all their cells, as well as animals which carry the transgene in some, but not all their cells, I. c., mosaic animals or chimeric. The transgene may be integrated as a single transgene or as multiple copies such as in concatamers, e. g., head-to-head tandems or head-to-tail tandems. The transgene may also be selectively introduced into and activated in a particular cell type by following, for example, the teaching of Lasko et al., (Lasko et al., Proc. Natl. Acad. Sci. USA 89: 6232-6236 (1992)). The regulatory sequences required for

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such a cell-type specific activation will depend upon the particular cell type of interest, and will be apparent to those of skill in the art. When it is desired that the polynucleotide transgene be integrated into the chromosomal site of the endogenous gene, gene targeting is preferred. Briefly, when such a technique is to be utilized, vectors containing some nucleotide sequences homologous to the endogenous gene are designed for the purpose of integrating, via homologous recombination with chromosomal sequences, into and disrupting the function of the nucleotide sequence of the endogenous gene. The transgene may also be selectively introduced into a particular cell type, thus inactivating the endogenous gene in only that cell type, by following, for example, the teaching of Gu et al. (Gu et al., Science 265: 103-106 (1994)). The regulatory sequences required for such a cell-type specific inactivation will depend upon the particular cell type of interest, and will be apparent to those of skill in the art.

Once transgenic animals have been generated, the expression of the recombinant gene may be assayed utilizing standard techniques. Initial screening may be accomplished by Southern blot analysis or PCR techniques to analyze animal tissues to verify that integration of the transgene has taken place. The level of mRNA expression of the transgene in the tissues of the transgenic animals may also be assessed using techniques which include, but are not limited to, Northern blot analysis of tissue samples obtained from the animal, in situ hybridization analysis, and reverse transcriptase-PCR (rt-PCR). Samples of transgenic gene-expressing tissue may also be evaluated immunocytochemically or immunohistochemically using antibodies specific for the transgene product.

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Once the founder animals are produced, they may be bred, inbred, outbred, or crossbred to produce colonies of the particular animal. Examples of such breeding strategies include, but are not limited to: outbreeding of founder animals with more than one integration site in order to establish separate lines; inbreeding of separate lines in order to produce compound transgenics that express the transgene at higher levels because of the effects of additive expression of each transgene; crossing of heterozygous transgenic animals to produce animals homozygous for a given integration site in order to both augment expression and eliminate the need for screening of animals by DNA analysis; crossing of separate homozygous lines to produce compound heterozygous or homozygous lines; and breeding to place the transgene on a distinct background that is appropriate for an experimental model of interest.

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Transgenic animals of the invention have uses which include, but are not limited to, animal model systems useful in elaborating the biological function of polypeptides of the present invention, studying conditions and/or disorders associated with aberrant expression, and in screening for compounds effective in ameliorating such conditions and/or disorders.

Example 14: Knock-Out Animals

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Endogenous gene expression can also be reduced by inactivating or"knocking out"the gene and/or its promoter using targeted homologous recombination. (E. g., see Smithies et al., Nature 317: 230-234 (1985); Thomas & Capecchi, Cell 51: 503512 (1987); Thompson et al., Cell 5: 313-321 (1989)), For example, a mutant, non-functional polynucleotide of the invention (or a completely unrelated DNA sequence) flanked by DNA homologous to the endogenous polynucleotide sequence (either the coding regions or regulatory regions of the gene) can be used, with or without a selectable marker and/or a negative selectable marker, to transfect cells that express polypeptides of the invention in vivo. In another embodiment, techniques known in the art are used to generate knockouts in cells that contain, but do not express the gene of interest. Insertion of the DNA construct, via targeted homologous recombination, results in inactivation of the targeted gene. Such approaches are particularly suited in research and agricultural fields where modifications to embryonic stem cells can be used to generate animal offspring with an inactive targeted gene (e.g., see Thomas & Capecchi 1987 and Thompson 1989, supra). However this approach can be routinely adapted for use in humans provided the recombinant DNA constructs are directly administered or targeted to the required site in vivo using appropriate viral vectors that will be apparent to those of skill in the art.

In further embodiments of the invention, cells that are genetically engineered to express the polypeptides of the invention, or alternatively, that are genetically engineered not to express the polypeptides of the invention (e. g., knockouts) are administered to a patient in vivo. Such cells may be obtained from the patient (I. e., animal, including human) or an MHC compatible donor and can include, but are not limited to fibroblasts, bone marrow cells, blood cells (e. g., lymphocytes), adipocytes, muscle cells, endothelial cells etc. The cells are genetically engineered in vitro using recombinant DNA techniques to introduce the coding sequence of polypeptides of the invention into the cells, or alternatively, to disrupt the coding sequence and/or endogenous regulatory sequence

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associated with the polypeptides of the invention, e. g., by transduction (using viral vectors, and preferably vectors that integrate the transgene into the cell genome) or transfection procedures, including, but not limited to, the use of plasmids, cosmids, YACs, naked DNA, electroporation, liposomes, etc.

The coding sequence of the polypeptides of the invention can be placed under the control of a strong constitutive or inducible promoter or promoter/enhancer to achieve expression, and preferably secretion, of the polypeptides of the invention. The engineered cells which express and preferably secrete the polypeptides of the invention can be introduced into the patient systemically, e. g., in the circulation, or intraperitoneally.

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Alternatively, the cells can be incorporated into a matrix and implanted in the body, e. g., genetically engineered fibroblasts can be implanted as part of a skin graft; genetically engineered endothelial cells can be implanted as part of a lymphatic or vascular graft. (See, for example, Anderson et al. U. S. Patent No. 5,399,349; and Mulligan & Wilson, U. S. Patent No. 5,460,959 each of which is incorporated by reference herein in its entirety).

When the cells to be administered are non-autologous or non-MHC compatible cells, they can be administered using well known techniques which prevent the development of a host immune response against the introduced cells. For example, the cells may be introduced in an encapsulated form which, while allowing for an exchange of components with the immediate extracellular environment, does not allow the introduced cells to be recognized by the host immune system.

Transgenic and "knock-out" animals of the invention have uses which include, but are not limited to, animal model systems useful in elaborating the biological function of polypeptides of the present invention, studying conditions and/or disorders associated with aberrant expression, and in screening for compounds effective in ameliorating such conditions and/or disorders

While preferred illustrative embodiments of the present invention are described, one skilled in the art will appreciate that the present invention can be practiced by other than the described embodiments, which are presented for purposes of illustration only and not by way of limitation. The present invention is limited only by the claims that follow.

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What is Claimed is:

- 1. An isolated nucleic acid molecule comprising:
- (a) a nucleic acid molecule comprising a nucleic acid sequence that encodes an
 amino acid sequence of SEQ ID NO: 109, 119, 120, 122, 123, 124, 125, 153, 166, 167,
 182, 183, 184, 185, 189 or 190;
 - (b) a nucleic acid molecule comprising a nucleic acid sequence of SEQ ID NO: 10, 20, 21, 24, 25, 26, 27, 58, 73, 74, 92, 93, 94, 95, 99 or 100;
 - (c) a nucleic acid molecule that selectively hybridizes to the nucleic acid molecule of (a) or (b); or
 - (d) a nucleic acid molecule having at least 90% sequence identity to the nucleic acid molecule of (a) or (b).
- The nucleic acid molecule according to claim 1, wherein the nucleic acid
 molecule is a cDNA.
 - The nucleic acid molecule according to claim 1, wherein the nucleic acid molecule is genomic DNA.
- 4. The nucleic acid molecule according to claim 1, wherein the nucleic acid molecule is a mammalian nucleic acid molecule.
 - The nucleic acid molecule according to claim 4, wherein the nucleic acid molecule is a human nucleic acid molecule.

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- A method for determining the presence of a colon specific nucleic acid (CSNA) in a sample, comprising the steps of:
- (a) contacting the sample with the nucleic acid molecule of SEQ ID NO: 9, 10, 11, 20, 21, 24, 25, 26, 27, 30, 31, 58, 72, 73, 74, 92, 93, 94, 95, 96, 99 or 100 under conditions in which the nucleic acid molecule will selectively hybridize to a colon specific nucleic acid: and
- (b) detecting hybridization of the nucleic acid molecule to a CSNA in the sample, wherein the detection of the hybridization indicates the presence of a CSNA in the sample.

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- A vector comprising the nucleic acid molecule of claim 1.
- 8. A host cell comprising the vector according to claim 7.

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- A method for producing a polypeptide encoded by the nucleic acid molecule according to claim 1, comprising the steps of:
- (a) providing a host cell comprising the nucleic acid molecule operably linked to one or more expression control sequences, and
- (b) incubating the host cell under conditions in which the polypeptide is produced.
 - 10. A polypeptide encoded by the nucleic acid molecule according to claim 1.
 - 11. An isolated polypeptide selected from the group consisting of:
- (a) a polypeptide comprising an amino acid sequence with at least 60% sequence identity to of SEQ ID NO: 109, 119, 120, 122, 123, 124, 125, 153, 166, 167, 182, 183, 184, 185, 189 or 190; or
- (b) a polypeptide comprising an amino acid sequence encoded by a nucleic acid molecule having at least 90% sequence identity to a nucleic acid molecule comprising a nucleic acid sequence of SEQ ID NO: 10, 20, 21, 24, 25, 26, 27, 58, 73, 74, 92, 93, 94, 95, 99 or 100.
 - 12. An antibody or fragment thereof that specifically binds to
- (a) a polypeptide comprising an amino acid sequence with at least 60%
 sequence identity to of SEQ ID NO: 108, 109, 110, 119, 120, 122, 123, 124, 125, 127,
 128, 153, 165, 166, 167, 182, 183, 184, 185, 186, 189 or 190; or
- (b) a polypeptide comprising an amino acid sequence encoded by a nucleic acid molecule having at least 90% sequence identity to a nucleic acid molecule comprising a nucleic acid sequence of SEQ ID NO: 9, 10, 11, 20, 21, 24, 25, 26, 27, 30, 31, 58, 72, 73, 74, 92, 93, 94, 95, 96, 99 or 100.

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PCT/HS02/27728

- 13. A method for determining the presence of a colon specific protein in a sample, comprising the steps of:
- (a) contacting the sample with a suitable reagent under conditions in which the reagent will selectively interact with the colon specific protein comprising an amino acid sequence with at least 60% sequence identity to of SEQ ID NO: 108, 109, 110, 119, 120, 122, 123, 124, 125, 127, 128, 153, 165, 166, 167, 182, 183, 184, 185, 186, 189 or 190; and
- (b) detecting the interaction of the reagent with a colon specific protein in the sample, wherein the detection of binding indicates the presence of a colon specific protein in the sample.

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- 14. A method for diagnosing or monitoring the presence and metastases of colon cancer in a patient, comprising the steps of:
- (a) determining an amount of the nucleic acid molecule of claim 1 or a polypeptide of claim 11 in a sample of a patient; and
- (b) comparing the amount of the determined nucleic acid molecule or the polypeptide in the sample of the patient to the amount of the colon specific marker in a normal control; wherein a difference in the amount of the nucleic acid molecule or the polypeptide in the sample compared to the amount of the nucleic acid molecule or the polypeptide in the normal control is associated with the presence of colon cancer.

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15. A kit for detecting a risk of cancer or presence of cancer in a patient, said kit comprising a means for determining the presence the nucleic acid molecule of claim 1 or a polypeptide of claim 11 in a sample of a patient.

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16. A method of treating a patient with colon cancer, comprising the step of administering a composition according to claim 11 or 12 to a patient in need thereof, wherein said administration induces an immune response against the colon cancer cell expressing the nucleic acid molecule or polypeptide.

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 A vaccine comprising the polypeptide or the nucleic acid encoding the polypeptide of claim 11.

1 SEQUENCE LISTING

<110> diaDexus, Inc. Sun, Yongming Liu, Chenghua Ghosh, Malavika <120> Compositions and Methods Relating to Colon Specific Genes and Proteins <130> DEX-0342 <150> US 60/316,259 <151> 2001-08-31 <160> 190 <170> PatentIn version 3.1 <210> 1 <211> 461 <212> DNA <213> Homo sapien <220> <221> misc_feature <222> (288)..(288) <223> n=a, c, g or t <220> <221> misc feature <222> (290)..(290) <223> n=a, c, q or t <220> <221> misc feature <222> (393)..(393) <223> n=a, c, g or t <400> 1 aqcaatatac taaqqqttac cttqccctca qtqqqcccaq aqtaqatqac acqatqtqqq 60 ataagaagtg gcccatgcac agaaggagag actccttttg tctgqagaca ctatggcgag 120 tttcttagac aaggtggtat ccgagatgag tttgcaaacg tgtgcacgat ttgatcatgc 180 agatgtggga ggaagagcat tccagatagg aggaatcacg tgaggatgag tatggtggca 240 ggaaagggag aaccatatat atataagaga ggaaaggaag tgttaganan ggtgttgcat 300 gcatgaaagg gaagaggagg aagaaatgaa gttggcaaga caaatctgaa cctgatctgg 360 gagaatettg actateteat gatttttqag aqntqqtqca caaaacatet qtettqttta 420 cttqctqqaa tqttqtqqcc tqtctqqttt qctttttaqa c 461

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Ala Arg Glu Asn Ala Lys Asp Lys Ser Arg Leu Asn Glu Gln Leu Ala 20 25

Gln Ile Pro Ile Ser Leu Met Pro Leu Ser Gln Leu Ile His Thr Val

Ser

<210> 104

<211> 40

<212> PRT <213> Homo sapien

<400> 104

Met Lys Ile Pro Tyr Leu Lys Gly Lys Tyr Ser Leu Ile Leu Asn Cys 5 10

75

Asn Ala Gly Lys Pro Asn Cys Phe Gly Ile Cys Asn Leu Asn Val Leu 25 Gln Asn Leu Val Leu Lys Phe Asp <210> 105 <211> 51 <212> PRT <213> Homo sapien <400> 105 Met Ile Asn Asp Pro Lys Leu Leu Tyr Leu Ser Asn Pro Cys Val Pro 5 10 Phe Leu Leu Phe Lys Lys Thr Ile Ser Pro Cys Arg Cys Leu Ser Leu Phe Cys Phe Cys Val Ser Val Leu Ser Tyr Ile Phe Ser Pro Pro Ile Cys Cys Phe <210> 106 <211> 45 <212> PRT <213> Homo sapien <400> 106 Met Gln Arg Tyr Thr Leu Ser Thr Ile Gln Tyr Val Phe Pro Ala Phe 5 10 15 Leu Gly Ile Tyr Ile Ala Val Ala Ile Phe Leu Ala Lys Ile Gly Ser 20 25 Tyr Tyr Ser Phe Pro Ile Leu Phe Phe Lys Val Asn Phe 35 40 <210> 107 <211> 51 <212> PRT <213> Homo sapien

<400> 107
Met His Asp Phe Leu Arg Met Ser Leu Pro Ser Asp Val Gly Ser Asp

76

Glu Thr Tyr Ser His Lys Val Met Ser Ser Thr Met Leu Ser Leu Lys 20 25 30

Ala Phe Glu Glu Leu Gly Gly Gln Val Cys Trp Gln Arg Pro Val Ile 35 40 45

Pro Ala Val

<210> 108 <211> 928 <212> PRT

<212> PRT <213> Homo sapien

<400> 108

Met Ala Glu Gly Lys Glu Lys Gln Val Thr Ser Tyr Met Asp Gly Ser 1 $$ 5 $$ 10 $$ 15

Arg Pro Tyr Asp Val Ser Met Thr Tyr Ile His Lys Ala Gly Gly Pro

Asp Gln Gln Glu Leu Val Met Leu Thr Cys Thr Val Pro Leu Asp Ser 35 40 45

Cys Cys His Leu Pro Gln Ala Arg Thr Asn Tyr Arg Lys Tyr Phe Arg 50 55 60

Ser Glu Ala Ala Phe Thr Leu Ala Asp Phe Ile Tyr Lys Ser Met Ile 65 70 75 80

Arg Val Asn Ser Ser Arg Leu Val Arg Val Thr Gln Val Glu Asn Glu 85 90 95

Glu Lys Leu Lys Glu Leu Glu Gln Phe Ser Ile Trp Asn Phe Phe Ser 100 105 110

Ser Phe Leu Lys Glu Lys Leu Asn Asp Thr Tyr Val Asn Val Gly Leu 115 \$120\$

Tyr Ser Thr Lys Thr Cys Leu Lys Val Glu Ile Ile Glu Lys Asp Thr 130 135 140

Lys Tyr Ser Val Ile Val Ile Arg Arg Ser Trp Asp Val Ile Arg Val 145 150 155 160

Asn Ser Ser Arg Leu Val Arg Val Thr Gln Val Glu Asn Glu Glu Lys

,															
									77						
				165					170					175	
Leu	Lys	Glu	Leu 180	Glu	Gln	Phe	Ser	Ile 185	Trp	Asn	Phe	Phe	Ser 190	Ser	Phe
Leu	Lys	Glu 195	Lys	Leu	Asn	Asp	Thr 200	Tyr	Val	Asn	Val	Gly 205	Leu	Tyr	Ser
Thr	Lys 210	Thr	Cys	Leu	Lys	Val 215	Glu	Ile	Ile	Glu	Lys 220	Asp	Thr	Lys	Tyr
Ser 225	Val	Ile	Val	Ile	Arg 230	Arg	Ser	Trp	Asp	Val 235	Ile	Arg	Val	Asn	Ser 240
Ser	Arg	Leu	Val	Arg 245	Val	Thr	Gln	Val	Glu 250	Asn	Glu	Glu	Lys	Leu 255	Lys
Glu	Leu	Glu	Gln 260	Phe	Ser	Ile	Trp	Asn 265	Phe	Phe	Ser	Ser	Phe 270	Leu	Lys
Glu	Lys	Leu 275	Asn	Азр	Thr	Tyr	Val 280	Asn	Gly	Ile	Pro	Trp 285	Thr	Lys	Val
Asp	Tyr 290	Phe	Asp	Asn	Gly	Ile 295	Ile	Cys	Lys	Leu	Ile 300	Glu	His	Asn	Gln
Arg 305	Gly	Ile	Leu	Ala	Met 310	Leu	Asp	Glu	Glu	Cys 315	Leu	Arg	Pro	Gly	Val 320
Val	Ser	Asp	Ser	Thr 325	Phe	Leu	Ala	Lys	Leu 330	Asn	Gln	Leu	Phe	Ser 335	Lys
His	Gly	His	Tyr 340	Glu	Ser	Lys	Val	Thr 345	Gln	Asn	Ala	Gln	Arg 350	Gln	Tyr
Asp	His	Thr 355	Met	Gly	Leu	Ser	Cys 360	Phe	Arg	Ile	Cys	His 365	Tyr	Ala	Gly
Lys	Val 370	Thr	Tyr	Asn	Val	Thr 375	Ser	Phe	Ile	Asp	Lys 380	Asn	Asn	Asp	Leu

Leu Arg Ser Leu Phe Pro Glu Gly Asn Pro Lys Gln Ala Ser Leu Lys 405 410 415

Leu Phe Arg Asp Leu Leu Gln Ala Met Trp Lys Ala Gln His Pro Leu

395

Arg Pro Pro Thr Ala Gly Ala Gln Phe Lys Ser Ser Val Ala Ile Leu 420 425 Met Lys Asn Leu Tyr Ser Lys Ser Pro Asn Tyr Ile Arg Cys Ile Lys Pro Asn Glu His Gln Gln Arg Gly Gln Phe Ser Ser Asp Leu Val Ala Thr Gln Ala Arg Tyr Leu Gly Leu Leu Glu Asn Val Arg Val Arg Arg 465 470 475 Ala Gly Tyr Ala His Arg Gln Gly Tyr Gly Pro Phe Leu Glu Arg Tyr 485 490 495 Arg Leu Leu Ser Arg Ser Thr Trp Pro His Trp Asn Gly Gly Asp Arg 500 505 Glu Gly Val Glu Lys Val Leu Gly Glu Leu Ser Met Ser Ser Gly Glu Leu Ala Phe Gly Lys Thr Lys Ile Phe Ile Arg Ser Pro Lys Thr Leu Phe Tyr Leu Glu Glu Gln Arg Arg Leu Arg Leu Gln Gln Leu Ala Thr 545 550 555 Leu Ile Gln Lys Ile Tyr Arg Gly Trp Arg Cys Arg Thr His Tyr Gln 570 Leu Met Arg Lys Ser Gln Ile Leu Ile Ser Ser Trp Phe Arg Gly Asn Met Ala Arq Lys Asn Tyr Arg Lys Tyr Phe Arg Ser Glu Ala Ala Leu 600 Thr Leu Ala Asp Phe Ile Tyr Lys Ser Met Val Gln Lys Phe Leu Leu Gly Leu Lys Asn Asn Leu Pro Ser Thr Asn Val Leu Asp Lys Thr Trp 625 630 635

Pro Ala Ala Pro Tyr Lys Cys Leu Ser Thr Ala Asn Gln Glu Leu Gln

650

885

Gln Leu Phe Tyr Gln Trp Lys Ala Thr Pro Val Pro Pro Ser Ser Gln 665 Cys Lys Arg Phe Arg Asp Gln Leu Ser Pro Lys Gln Val Glu Ile Leu 680 Arg Glu Lys Leu Cys Ala Ser Glu Leu Phe Lys Gly Lys Lys Ala Ser Tyr Pro Gln Ser Val Pro Ile Pro Phe Cys Gly Asp Tyr Ile Gly Leu 710 Gln Gly Asn Pro Lys Leu Gln Lys Leu Lys Gly Gly Glu Glu Gly Pro Val Leu Met Ala Glu Ala Val Lys Lys Val Asn Arg Gly Asn Gly Lys Thr Ser Ser Arg Ile Leu Leu Thr Lvs Glv His Val Ile Leu Thr 760 Asp Thr Lys Lys Ser Gln Ala Lys Ile Val Ile Gly Leu Asp Asn Val Ala Gly Val Ser Val Thr Ser Leu Lys Asp Gly Leu Phe Ser Leu His 790 795 Leu Ser Glu Met Ser Ser Val Gly Ser Lys Gly Asp Phe Leu Leu Val 805 810 815 Ser Glu His Val Ile Glu Leu Leu Thr Lys Met Tyr Arg Ala Val Leu 820 825 Asp Ala Thr Gln Arg Gln Leu Thr Val Thr Val Thr Glu Lys Phe Ser 835 840 Val Arg Phe Lys Glu Asn Ser Val Ala Val Lys Val Val Gln Gly Pro 850 Ala Gly Gly Asp Asn Ser Lys Leu Arg Tyr Lys Lys Lys Gly Ser His 865 Cys Leu Glu Val Thr Val Gln Gln Leu Thr Ala Gly Tyr His Ala Gly

R۸

Gln Gly Glu Leu Ile Asn Phe Ser Ser Cys Leu Gln Ile Asn Leu Leu 905

Ser Glu His Lys Pro Arg Ala Ser Gly Thr Pro Cys Phe Glu Leu Arg 920

<210> 109 <211> 351 <212> PRT

<213> Homo sapien

<400> 109

Met Glu Pro Thr Glu Pro Met Glu Pro Thr Glu Pro Met Glu Pro Thr 10

Glu Pro Met Glu Pro Thr Glu Pro Met Glu Pro Thr Glu Pro Met Glu 25

Pro Ala Arg Ser Ala His Arg Gly Gly Glu Ala Leu Leu Arg Glu Leu 40

Glu Val Leu Val Gln Asp Val Val Arg Thr Ser Ser Trp Trp Glu Arg

His Gly Val Asp Cys Ala Ile Leu Ala Leu Ser Leu Phe Ala Leu Pro 65 70 75

Ala Gly Phe Leu Cys Leu Arg Trp Glu Asn Ala Leu Val Phe Ala Ser 90 85

Gly Ile Thr Ile Leu Gly Val Cys His Tyr Thr Leu Thr Val Lys Gly 100 105

Ser His Leu Ala Thr His Gly Ala Leu Thr Glu Ser Lys Arg Trp Ser 115 120

Lys Ile Trp Leu Leu Phe Phe Val Glu Val Cys Thr Ala Phe Thr Ala 130 135 140

Glu His Ala Thr His Gly His Val Lys Met His His Ala Tyr Thr Asn 145

Val Val Gly Leu Gly Asp Ser Ser Thr Trp Arg Leu Pro Cys Leu Asn 165 170

Arg Tyr Val Tyr Met Phe Leu Ala Pro Phe Leu Leu Pro Ile Ala Thr 180 185 190

81

Pro Leu Val Ala Val Glu Arg Leu Arg Lys Val Glu Leu Gly Thr Ala 195 200 Leu Arg Thr Leu Ala Leu Ile Ser Leu Gly Leu Tyr Ser His Tyr Trp Leu Leu Leu Asn Val Ser Gly Phe Lys Asn Pro Ser Ser Ala Leu Gly 225 235 Cys Met Phe Leu Thr Arg Ser Leu Leu Ala His Pro Tyr Leu His Val 245 250 255 Asn Ile Phe Gln His Ile Gly Leu Pro Met Phe Ser Arg Asp Asn Lys 260 Pro Arg Arg Ile His Met Met Ser Leu Gly Val Leu Asn Leu Ala Arg 280 Leu Pro Val Leu Asp Trp Ala Phe Gly His Ser Ile Ile Ser Cys His 290 295 300 Val Glu His His Leu Phe Pro Arg Leu Ser Asp Asn Met Cys Leu Lys 305 315 Val Val Glu Gly Trp Ala Gly Gly Ala Gly Ile Lys Gly Leu Leu Glu 325 330 Asp Gly Lys Glu Asp Ser Tyr Gly Leu Gly Ala Leu Leu Thr Leu 345 <210> 110 <211> 715 <212> PRT <213> Homo sapien <400> 110 Met Arg Gln Ser Leu Leu Phe Leu Thr Ser Val Val Pro Phe Val Leu 5 Ala Pro Arg Pro Pro Asp Asp Pro Gly Phe Gly Pro His Gln Arg Leu 20

Glu Lys Leu Asp Ser Leu Leu Ser Asp Tyr Asp Ile Leu Ser Leu Ser

40

									82						
Asn	Ile 50	Gln	Gln	His	Ser	Val 55	Arg	Lys	Arg	Asp	Leu 60	Gln	Thr	Ser	Thr
His 65	Val	Glu	Thr	Leu	Leu 70	Thr	Phe	Ser	Ala	Leu 75	Lys	Arg	His	Phe	Lys 80
Leu	Tyr	Leu	Thr	Ser 85	Ser	Thr	Glu	Arg	Phe 90	Ser	Gln	Asn	Phe	Lув 95	Val
Val	Val	Val	Asp 100	Gly	Lys	Asn	Glu	Ser 105	Glu	Tyr	Thr	Val	Lys 110	Trp	Gln
Asp	Phe	Phe 115	Thr	Gly	His	Val	Val 120	Gly	Glu	Pro	Asp	Ser 125	Arg	Val	Leu
Ala	His 130	Ile	Arg	Asp	Asp	Asp 135	Val	Ile	Ile	Arg	Ile 140	Asn	Thr	Asp	Gly
Ala 145	Glu	Tyr	Asn	Ile	Glu 150	Pro	Leu	Trp	Arg	Phe 155	Val	Asn	Asp	Thr	Lys 160
Asp	Lys	Arg	Met	Leu 165	Val	туг	Lys	ser	Glu 170	Asp	Ile	Lys	Asn	Val 175	Ser
Arg	Leu	Gln	Ser 180	Pro	Lys	Val	Cys	Gly 185	Tyr	Leu	Lys	Val	Asp 190	Asn	Glu
Glu	Leu	Leu 195	Pro	Lys	Gly	Leu	Val 200	Asp	Arg	Glu	Pro	Pro 205	Glu	Glu	Leu
Val	His 210	Arg	Val	Lys	Arg	Arg 215	Ala	Asp	Pro	Asp	Pro 220	Met	Lys	Asn	Thr
Cys 225	Lys	Leu	Leu	Val	Val 230	Ala	Asp	His	Arg	Phe 235	туг	Arg	Tyr	Met	Gly 240
Arg	Gly	Glu	Glu	Ser 245	Thr	Thr	Thr	Asn	Tyr 250	Leu	Ile	Glu	Leu	11e 255	Asp
Arg	Val	Asp	Asp 260	Ile	Tyr	Arg	Asn	Thr 265	Ser	Trp	Asp	Asn	Ala 270	Gly	Phe
Lys	Gly	Tyr 275	Gly	Ile	Gln	Ile	Glu 280	Gln	Ile	Arg	Ile	Leu 285	Lys	Ser	Pro

Gln Glu Val Lys Pro Gly Glu Lys His Tyr Asn Met Ala Lys Ser Tyr

	290					295					300				
Pro 3 0 5	Asn	Glu	Glu	Lys	Asp 310	Ala	Trp	Asp	Val	Lys 315	Met	Leu	Leu	Glu	Gln 320
Phe	Ser	Phe	Asp	Ile 325	Ala	Glu	Glu	Ala	Ser 330	Lys	Val	Cys	Leu	Ala 335	His
Leu	Phe	Thr	Tyr 340	Gln	Asp	Phe	Asp	Met 345	Gly	Thr	Leu	Gly	Leu 350	Ala	Tyr
Val	Gly	Ser 355	Pro	Arg	Ala	Asn	Ser 360	His	Gly	Gly	Val	Cys 365	Pro	Lys	Ala
Tyr	Tyr 370	Ser	Pro	Val	Gly	Lys 375	Lys	Asn	Ile	Tyr	Leu 380	Asn	Ser	Gly	Leu
Thr 385	Ser	Thr	Lys	Asn	Tyr 390	Gly	Lys	Thr	Ile	Leu 395	Thr	Lys	Glu	Ala	Asp 400
Leu	Val	Thr	Thr	His 405	Glu	Leu	Gly	His	Asn 410	Phe	Gly	Ala	Glu	His 415	Asp
Pro	Asp	Gly	Leu 420	Ala	Glu	Cys	Ala	Pro 425	Asn	Glu	Asp	Gln	Gly 430	Gly	Lys
Tyr	Val	Met 435	Tyr	Pro	Ile	Ala	Val 440	Ser	Gly	Авр	His	Glu 445	Asn	Asn	Lys
Met	Phe 450	Ser	Asn	Cys	Ser	Lys 455	Gln	Ser	Ile	Tyr	Lys 460	Thr	Ile	Glu	Ser
Lys 465	Ala	Gln	Glu	Сув	Phe 470	Gln	Glu	Arg	Ser	Asn 475	Lys	Val	Cys	Gly	Asn 480
Ser	Arg	Val	Asp	Glu 485	Gly	Glu	Glu	Cys	Asp 490	Pro	Gly	Ile	Met	Tyr 495	Leu
Asn	Asn	Asp	Thr 500	Cys	Cys	Asn	Ser	Asp 505	Сув	Thr	Leu	Lys	Glu 510	Gly	Val
Gln	Сув	Ser 515	Asp	Arg	Asn	Ser	Pro 520	Сув	Сув	Lys	Asn	Cys 525	Gln	Phe	Glu
Thr	Ala 530	Gln	Lys	Lys	Сув	Gln 535	Glu	Ala	Ile	Asn	Ala 540	Thr	Сув	Lys	Gly

84

Val Ser Tyr Cys Thr Gly Asn Ser Ser Glu Cys Pro Pro Pro Gly Asn Ala Glu Asp Asp Thr Val Cys Leu Asp Leu Gly Lys Cys Lys Asp Gly Lys Cys Ile Pro Phe Cys Glu Arg Glu Gln Gln Leu Glu Ser Cys Ala Cys Asn Glu Thr Asp Asn Ser Cys Lys Val Cys Cys Arg Asp Leu Ser 595 600 605 Gly Arg Cys Val Pro Tyr Val Asp Ala Glu Gln Lys Asn Leu Phe Leu 610 615 Arg Lys Gly Lys Pro Cys Thr Val Gly Phe Cys Asp Met Asn Gly Lys 630 635 Cys Glu Lys Arg Val Gln Asp Val Ile Glu Arg Phe Trp Asp Phe Ile 645 650 Asp Gln Leu Ser Ile Asn Thr Phe Gly Lys Phe Leu Ala Asp Asn Ile Val Gly Ser Val Leu Val Phe Ser Leu Ile Phe Trp Ile Pro Phe Ser 680 Ile Leu Val His Cys Val Ile Arg Asn Trp Ile Asn Ser Met Asn Leu 695 Cys Leu Cys Phe Thr Pro Val Thr Ser Lys Cys 710 <210> 111 <211> 65 <212> PRT <213> Homo sapien <400> 111 Met Ala Ser Gly Gly Val Leu His Val Met Ser Gly Arg Lys Ser Gly

Val Phe Leu Arg Gln Cys Val Phe Met Trp Ser Lys Gln Ser Lys Pro 20 25

85

Val Ser Glu Ser Asn Pro Ser Met Thr Met Phe Pro His Leu Cys His 40

Thr Leu Cys Glu Glu Leu Cys Pro His Phe Ser Leu Phe Asn Asn Leu 55 60

Met 65

<210> 112 <211> 22 <212> PRT <213> Homo sapien

<400> 112

Met Asp His Tyr Ile Tyr Pro Val Asn Phe Pro Gly Ser Asn Cys Gly

Tyr Pro Asn Val Phe Glu 20

<210> 113

<211> 21 <212> PRT <213> Homo sapien

<400> 113

Met Lys Phe Gln Leu Phe Ser Met His Lys Asn Arg Tyr Tyr Asp Ile

Val His Tyr Thr Met 20

<210> 114 <211> 41 <212> PRT

<213> Homo sapien

<400> 114

Met Ala Lys Leu Val Thr Thr Ala Arg Ser Gly Ser Glu Arg Asp Asp

Lys Glu Gly Glu Phe Lys Glu Pro Gln Thr Pro Gly Ile Phe Cys Ala

Arg Ala Asn Asp Thr Glu Ser Ile Pro 35

86

<210> 115 <211> 42 <212> PRT

<213> Homo sapien

<220>

<221> MISC_FEATURE

<222> (42)..(42) <223> x=any amino acid

<400> 115

Met Glu Lys Thr Leu Ala Trp Leu Ser Lys Asp Met Gly Ala Asn Ser

Arg Leu Ala Leu Pro Ile Thr Tyr Cys Ala Gly Leu Thr Gln Ser Leu

Pro Leu Thr Arg Ser Gln Phe Leu His Xaa

<210> 116 <211> 28

<212> PRT <213> Homo sapien

<400> 116

Met Glu Thr His Leu Leu Met Arg Lys Gln Phe Thr Thr Cys Ser Ile

Glu His Ser Tvr Leu Glu Phe Asn Thr His Leu Tvr 20

<210> 117

<211> 97

<212> PRT

<213> Homo sapien

<400> 117

Met Pro Leu Ala Val Thr Gly Thr Cys His Ala Cys Ser Phe Ile Gly 5 10

His Cys Thr Cys Leu Leu Phe Ala Phe Lys Ala Leu Pro Leu Asp Ile 20

Arg Ile Ala Ser Phe Phe Ala Ser Phe Arg Phe Leu Thr Ile Cys His

Leu Leu Gly Glu Ala Phe Tyr Asp His Leu Thr Cys Asn Ser Ser Ser

87

Thr Pro Tyr Tyr Leu Ser Pro Ser Ser Val Leu Phe Phe Phe Thr Val 70

Tyr Phe Trp Leu Ile Val Ser Leu Pro Gln Asp Ile Val Ile Ser Gly

Gly

<210> 118 <211> 137 <212> PRT <213> Homo sapien

<400> 118

Met Cys Leu Gly Ser Gly Ile Thr Trp Leu Gly Pro Gln Ile Phe Ser

Ser Ala Trp Lys Arq Phe Thr Ser Ser Ala Ser Ser Arq Cys Gly Ser

Arg Gly Ile Asp Gly Leu Leu Thr Ser Thr Phe Ser Phe Pro Ala His

Leu Ala Leu Leu Gly His Val Ser Pro Val His Leu Gln Glu Thr 50 55 60

Ser Val Asp Ala Pro Cys Leu Leu Thr Leu Ser Pro Ala His Thr Glu

Leu Val Leu Arg Gly Asn Leu Cys Leu Cys Cys Cys Leu Cys Leu Glu 85 90 95

Arg Pro Cys Pro Thr Pro Ser His Ser Cys Leu Ser Val Ile Phe Pro 100 105

Met Ser Val Leu Arg Glu Pro Phe Leu Ala Thr Pro Ser Lys Gly Val 115

Leu Gly Gln Val Trp Trp Pro Thr Pro 130 135

<210> 119 <211> 51

<212> PRT <213> Homo sapien

88

<400> 119

Met Ser Val Leu Arg Glu Pro Phe Leu Ala Thr Pro Ser Lys Gly Val

Leu Gly Gln Val Trp Trp Pro Thr Pro Val Ile Ser Glu Leu Trp Glu

Thr Glu Val Gly Arg Ser Leu Glu Ala Arg Gly Ser Arg Pro Thr Trp 35 40 45

Ala Thr Tyr 50

<210> 120

<211> 89 <212> PRT

<213> Homo sapien

<400> 120

Met Ser Leu Cys Leu Ala Cys Thr Leu Cys Leu Gly Cys Ser Leu His 1 $$ 5 $$ 10 $$ 15

Pro Ala Leu Pro Gly Trp Ala Ser Asp Ser Asn Leu Leu Gly Leu Pro $20 \hspace{1cm} 25 \hspace{1cm} 30 \hspace{1cm}$

Pro His Leu Cys Asp Ser Gly Ile Ile Pro Lys Ala Val Val Arg Ile 35 40 45

Leu Gln Glu Asn Ala Trp Lys Val Leu Gly Thr Met Leu Ser Pro Tyr 50 55 60

Asp Thr His Ser Cys Val Leu Leu Ser Leu Thr Tyr Cys Phe Ser Thr 65 70 75 80

Thr Thr Thr Ile Arg Ile Leu Lys Val

<210> 121

<211> 64 <212> PRT

<213> Homo sapien

<400> 121

Met Pro Ala Val Thr Ile Thr Ile Met Tyr Phe Cys Cys Cys His Thr 1 5 10 15

89 Lys Met Asn Asn Asn Ile Leu Ser His Leu Lys Pro Lys Arg Arg Asn 20 25 Gln Trp Glu Gly Cys Leu Gln Pro Ala His Gln His Arg Lys Gly Ser 40 Pro Ala Ser Tyr Pro Asn Ser Gln Arg Pro Asn Pro Arg Leu Leu His <210> 122 <211> 34 <212> PRT <213> Homo sapien <400> 122 Met Arg Glu His Asn Asn His Asn Asp Asn Glu Cys Ser Val Val Lys Leu Thr Gly Thr Leu Leu Phe Leu Leu Ser Val Gln Pro Asn Ala Ser 25 20 Ala Asp <210> 123 <211> 151 <212> PRT <213> Homo sapien <400> 123 Met Tyr Gly Cys Tyr Thr Pro Thr Ala Tyr Ser Thr Arg Ser Ala Pro Glu Glu Asp Trp Val Lys Leu Cys Lys Phe Gly Phe Pro Gly Asn Ala Leu His Tyr Ser Ala Pro Asp Leu Pro Thr Thr Pro Val Gly Thr Arg Ser Ser Thr His Leu Ala Glu Leu Met Thr Ala Trp Ala Gln Arg Ser 55

Ala His Cys Ala Asn Thr Arg Thr Gly Ile Ala Pro Leu Pro Glu Pro

Pro His Arg Ala Pro Phe Lys Glu Leu Ala Thr Pro Leu Thr Cys Lys

75

70

90

Gln Pro Pro Thr Leu Lys Leu Ile Arg Thr Arg Val Phe His Pro Lys 105

Gly Leu Cys Cys Gly Arg Cys Ser Asp Pro Arg Arg Gly Arg Glu Val 120

Pro Lys Ala Thr Ala Arg Gly Trp Gly Thr Pro Leu Leu Thr Leu Val

Leu Asp Phe Glu Gly Pro Asn

<210> 124 <211> 59

<212> PRT

<213> Homo sapien

<400> 124

Met Asn Cys Leu Trp Ile Leu Leu Ser Ile Ser Leu Val Pro Phe Leu

Gln Leu Tyr Gly Thr Leu Ser Ser Cys Thr Pro Glu Ala Pro Gln Leu

Gly Lys Val Ser Gln Arg Tyr Gln Glu Tyr Met Leu Arg Gly His Phe

Lys Val Phe His Arq Arq Leu Cys Leu Glv Lys

<210> 125 <211> 83

<212> PRT

<213> Homo sapien

<400> 125

Met Asp Leu Cys Ile Ser Arg Ser Cys Ser Leu Leu Ser Val Gly Val

Leu Thr Arg Lys Ser Leu Ala Val Gln His Arg Cys Thr Ala Thr Leu

Gly Ala Val Glu Trp Ser Ala Gly Ser Gln Ser Asn Cys Phe Pro Leu 40

Asp Leu Gly Leu Val Leu Phe Ser Lys Tyr Gln Ser Tyr Leu Lys Met

91

Lys Ala Cys Lys Pro Leu Leu Lys Ser Ile Glu Ser Glu Arg Glu Arg 70 75

Leu Phe Cys

<210> 126 <211> 34

<212> PRT <213> Homo sapien

<400> 126

Met Lys Lys Ile Phe Lys Phe Asn Lys Ile Ile Ile Tyr Cys Pro Lys

Ile Ile Ile Lys Ser Leu Leu Val Glu Leu Ser Gln Asn Lys Gly Asn 20 25

Ser Phe

<210> 127

<211> 862 <212> PRT <213> Homo sapien

<400> 127

Met Met Pro Trp Ala Leu Gln Lys Lys Arg Glu Ile His Met Ala Lys

Ala His Arg Arg Arg Ala Ala Arg Ser Ala Leu Pro Met Arg Leu Thr 25

Ser Cys Ile Phe Arg Arg Pro Val Thr Arg Ile Arg Ser His Pro Asp

Asn Gln Val Arg Arg Arg Lys Gly Asp Glu His Leu Glu Lys Pro Gln

Gln Leu Cys Ala Tyr Arg Arg Leu Gln Ala Leu Gln Pro Cys Ser Ser 70

Gln Gly Glu Gly Ser Ser Pro Leu His Leu Glu Ser Val Leu Ser Ile 85 90

Leu Ala Pro Gly Thr Ala Ser Glu Ser Leu Asp Arg Ala Gly Ala Glu

			100					105					110		
Arg	Val	Arg 115	Ser	Pro	Leu	Glu	Pro 120	Thr	Pro	Gly	Arg	Phe 125	Pro	Ala	Val
Ala	Gly 130	Gly	Pro	Thr	Pro	Gly 135	Met	Gly	Cys	Gln	Leu 140	Pro	Pro	Pro	Leu
Ser 145	Gly	Gln	Leu	Val	Thr 150	Pro	Ala	Авр	Ile	Arg 155	Arg	Gln	Ala	Arg	Arg 160
Val	Lys	Lys	Ala	Arg 165	Glu	Arg	Leu	Ala	Lys 170	Ala	Leu	Gln	Ala	Asp 175	Arg
Leu	Ala	Arg	Arg 180	Ala	Glu	Met	Leu	Thr 185	Gly	Arg	Gln	Thr	Ile 190	Pro	Lys
туг	Asn	Cys 195	Gln	Thr	Gln	Asn	Lys 200	Thr	Val	Ala	Val	Met 205	Pro	Gly	Thr
Thr	Ser 210	Val	Ser	Ser	Thr	Lys 215	Ile	Gly	Ala	Leu	Leu 220	Gly	Cys	Tyr	Arg
Ser 225	Pro	Arg	Gly	Ala	Val 230		Arg	Ala	Leu	Arg 235	Asp	Lys	Gly	Gly	Ala 240
Ser	Cys	Phe	Pro	Ala 245	Ala	Leu	Leu	Leu	Ser 250	Glu	Ser	Gln	Asp	Gly 255	Gly
Cys	Arg	Pro	Ala 260	Pro	Asn	Cys	Arg	Ala 265	Met	Leu	Trp	Phe	Pro 270	Trp	Ile
Ala	Val	Gly 275	Trp	Cys	Leu	Leu	Gly 280	Val	Glu	Val	Asp	Ala 285	Leu	Trp	Val
Thr	Val 290	Tyr	Ala	Trp	Gly	Pro 295	Gly	Arg	Ser	Gln	Ala 300	Ala	Ser	Gln	Arg
Glu 305	Gly	His	Thr	Glu	Gly 310	Gly	Glu	Leu	Val	Asn 315	Glu	Leu	Leu	Lys	Ser 320
Trp	Leu	Lys	Gly	Leu 325	Val	Thr	Phe	Glu	Asp 330	Val	Ala	Val	Glu	Phe 335	Thr
Gln	Glu	Glu	Trp 340	Ala	Leu	Leu	Asp	Pro 345	Ala	Gln	Arg	Thr	Leu 350	Tyr	Arg

Asp Val Met Leu Glu Asn Cys Arg Asn Leu Ala Ser Leu Gly Asn Gln 355 360 Val Asp Lys Pro Arg Leu Ile Ser Gln Leu Glu Gln Glu Asp Lys Val 375 Met Thr Glu Glu Arg Gly Ile Leu Ser Gly Thr Cys Pro Asp Val Glu 395 390 Asn Pro Phe Lvs Ala Lvs Glv Leu Thr Pro Lvs Leu His Val Phe Arg 405 410 Lys Glu Gln Ser Arg Asn Met Lys Met Glu Arg Asn His Leu Gly Ala 420 425 Thr Leu Asn Glu Cys Asn Gln Cys Phe Lys Val Phe Ser Thr Lys Ser 440 Ser Leu Thr Arq His Arq Lys Ile His Thr Gly Glu Arg Pro Tyr Gly 450 455 Cys Ser Glu Cys Gly Lys Ser Tyr Ser Ser Arg Ser Tyr Leu Ala Val His Lys Arg Ile His Asn Gly Glu Lys Pro Tyr Glu Cys Asn Asp Cys 485 490 Gly Lys Thr Phe Ser Ser Arg Ser Tyr Leu Thr Val His Lys Arg Ile His Asn Gly Glu Lys Pro Tyr Glu Cys Ser Asp Cys Gly Lys Thr Phe 520 Ser Asn Ser Ser Tyr Leu Arg Pro His Leu Arg Ile His Thr Gly Glu 535 Lys Pro Tyr Lys Cys Asn Gln Cys Phe Arg Glu Phe Arg Thr Gln Ser Ile Phe Thr Arg His Lys Arg Val His Thr Gly Glu Gly His Tyr Val Cys Asn Gln Cys Gly Lys Ala Phe Gly Thr Arg Ser Ser Leu Ser Ser 580 585 590

His Tyr Ser Ile His Thr Gly Glu Tyr Pro Tyr Glu Cys His Asp Cys 595 Gly Arg Thr Phe Arg Arg Arg Ser Asn Leu Thr Gln His Ile Arg Thr 615 His Thr Gly Glu Lys Pro Tyr Thr Cys Asn Glu Cys Gly Lys Ser Phe Thr Asn Ser Phe Ser Leu Thr Ile His Arg Arg Ile His Asn Glv Glu 645 650 Lys Ser Tyr Glu Cys Ser Asp Cys Gly Lys Ser Phe Asn Val Leu Ser Ser Val Lys Lys His Met Arg Thr His Thr Gly Lys Lys Pro Tyr Glu 680 Cvs Asn Tyr Cys Gly Lys Ser Phe Thr Thr Ser Thr Thr Ser Thr Thr 695 Ala Ser Ala Thr Ser Thr Thr Ala Ser Trp Ile His His Ser Asn His 715 His Ile His Tyr Ser Ile His Tyr Arg Ile His Tyr Ser Asp Ser Asn 725 730 Thr Val Ala Ser His Ser Trp Gly Tyr Ala Ser Gly Asn Gly Arg Val 740 745 750 Glu Glu Gly Gly Arg Asp Pro Arg Val Val Ala Ser Thr Gly Asn Asp 755 760 Pro Glu Ala Ser Asp Asn Asp Glu Asp Asn Asp Gly Asp Asp Lys Asn 770 775 780 Gly His Asp Gly Asp Asp Ser Asp Gly Asn Glu Gly Asp Gly Asp Glu Met Val Met Met Met Ile Leu Thr Met Met Val Met Thr Val Ile 805 Met Met Met Val Met Leu Met Thr Val Met Glu Met Ala Asn Leu Thr 820 825 830

95

Thr Pro Tyr Cys Gly Thr Leu Gly Gln Ser Leu Glu Glu Cys Glu Glu 835 840 845

Phe Arg Arg Asn Phe His Ala Ala Ser Gly Lys Leu Pro Gly 850 860

<210> 128

<211> 187

<212> PRT <213> Homo sapien

<400> 128

Met Met Pro Trp Ala Leu Gln Lys Lys Arg Glu Ile His Met Ala Lys 1 $$ 5 $$ 10 $$ 15

Ala His Arg Arg Arg Ala Ala Arg Ser Ala Leu Pro Met Arg Leu Thr 20 25 30

Ser Cys Ile Phe Arg Arg Pro Val Thr Arg Ile Arg Ser His Pro Asp $35 \ \ \, 40 \ \ \, 45$

Asn Gln Val Arg Arg Arg Lys Gly Asp Glu His Leu Glu Lys Pro Gln 50 60

Gln Leu Cys Ala Tyr Arg Arg Leu Gln Ala Leu Gln Pro Cys Ser Ser 65 70 75 80

Gln Gly Glu Gly Ser Ser Pro Leu His Leu Glu Ser Val Leu Ser Ile 85 90 95

Leu Ala Pro Gly Thr Ala Gly Glu Ser Leu Asp Arg Ala Gly Ala Glu 100 105 110

Arg Val Arg Ser Pro Leu Glu Pro Thr Pro Gly Arg Phe Pro Ala Val

Ala Gly Gly Pro Thr Pro Gly Met Gly Cys Gln Leu Pro Pro Pro Leu 130 135 140

Ser Gly Gln Leu Val Thr Pro Ala Asp Ile Arg Arg Gln Ala Arg Arg 145 150155155

Val Lys Lys Ala Arg Glu Arg Leu Ala Lys Ala Leu Gln Ala Asp Arg 165 170 175

Leu Ala Arg Gln Ala Glu Met Leu Thr Cys Arg

96

<210> 129 <211> 11 <212> PRT <213> Homo sapien

<400> 129

Met Arg Lys Leu His Phe Glu Lys Tyr Arg Tyr

<210> 130 <211> 109 <212> PRT <213> Homo sapien

<400> 130

Met Ala Gly Asn Leu Thr Gln Pro Ala Gly Gln His Thr Thr Pro Glu 10

Pro Ser His Ala Gly Gly Cys Pro Gln Val Pro Lys Gly Leu Glu Gln

Ala Thr Leu Gly Asp Cys Leu His Pro Ile Leu Gln Arg Ala Val Cys

Gln Arg Val Pro Ala Ala Ser Gln Thr Ala Ile Ile His Gly Glu Met

Leu Ala Thr Val Pro Ser Thr Lys Val Tyr Ser Gln Cys Ile Ser Leu 65 70 75

Arg Leu Tyr Gly Gln Arg Val Gly Tyr His Leu Lys Gly Gln Lys Ala

Glu Pro Leu Ser Cys Trp Gly Ser Gly His Gln Leu Val 100 105

<210> 131 <211> 127

<212> PRT

<213> Homo sapien

<400> 131

Met Leu Ile Ile Leu Ala Leu Trp Asp Ala Glu Val Glu Gly Pro Leu

Glu Ala Ser Leu Gln Ala Pro Glu Glu Thr Ser Ser Met Ile Pro Cys 25

Leu Arg Gly Arg Phe Leu Arg Thr Leu Pro Trp Gly Leu Lys Val Ala

Thr Pro Pro Gly Thr Tyr Ala Pro Gly Ala Glu Phe Gly Leu Pro Lys 50 60

Ser Val Cys Val Arg Leu Cys Leu Cys Ala Cys Leu His Val Cys Val 65 70 75 80

Phe Ala Cys Met Ser Val Cys Leu Cys Gly Leu Cys Val Cys Val Tyr 85 90 95

Thr Ser Val Cys Leu Ser Leu Cys Ile Phe Ala Cys Val Ser Met Cys

Leu Cys Ala Cys Leu Cys Val Tyr Met Ser Val Ser Leu Tyr Leu 115 120 125

<210> 132

<211> 78 <212> PRT

<213> Homo sapien

<400> 132

Met His Ile Tyr Val Ile Ser Thr Asn Ser Pro Phe Val Lys Thr Asn 1 $$ 10 $$ 15

His Met Ala Lys Pro Ile Val Arg Gly Lys Tyr Thr Val Phe Arg $20 \ 25 \ 25 \ 30$

Ala Asn Tyr Arg Phe Thr Gln Gln Lys Ala Trp Ile Gln Glu Arg Val 35 40 45

Lys Gly Ser Ile Ile His Ser Thr Asn Arg Arg Ile Ser Arg Leu Arg 50 55

Pro Lys Asn Arg Cys Gln Gly His Val His His Thr Asp Ile 65 70 75

<210> 133

<211> 56 <212> PRT

<213> Homo sapien

<400> 133

Met Gly Gly Arg Glu Asp Arg Pro Gly Val Trp Asp Val Thr Ser Ala

15

98 1 5 10

His Thr Gln Gly Arg Pro Ala Gln Gly Trp Gly Leu Leu Cys Leu

Met Cys Ala Gly Ser Asp Thr Gly Leu Val Trp Gly Arg Leu Arg Thr

Val Lys Met Lys Asn Lys Arg Lys

<210> 134 <211> 53

<212> PRT

<213> Homo sapien

<400> 134

Met Val Pro His Pro Gly Phe Ser Gly Met Thr Trp Gly Thr Val Lys

Asn Thr Asp Ala Trp Val Pro Pro Pro Glu Ile Leu Ile Glv Leu Lvs 25

Gly Gln Val Gln Glu Ser Gly Val Leu Glu Val Pro Gln Val Ile Leu 40

Ile Gly Ser Gln Gly

<210> 135 <211> 47

<212> PRT

<213> Homo sapien

<400> 135

Met Asn Pro Ile Ala Phe Phe Asp Ser Gln Asn Ser Leu Gln Asn Arg

Leu Ser Ala Phe Gly Leu Ile Phe Thr Asn His Phe Pro Cys Ser Asn

Met Tyr Leu Arg Val Leu Thr Leu Leu Ser Val Thr Met Phe Tyr 40

<210> 136 <211> 162 <212> PRT

<213> Homo sapien

99

<400> 136

Met Val Arg Arg Phe Leu Val Thr Leu Arg Ile Arg Arg Ala Cys Gly

Pro Pro Arg Val Arg Val Phe Val Val His Ile Pro Arg Leu Thr Gly

Glu Trp Ala Ala Pro Gly Ala Pro Ala Ala Val Ala Leu Val Leu Met 35 40 45

Leu Leu Arg Ser Gln Arg Leu Gly Gln Gln Pro Leu Pro Arg Arg Pro 50 55 60

Gly Arg Lys Gly Pro Arg Lys Val Arg Gly Ala Phe Gly Thr Cys Phe 65 70 75 80

Val Cys Thr Val Glu Pro Gly Pro Pro Phe Thr Gln Pro Leu Asn Arg 85 90 95

Gly Lys Leu Arg Arg Thr Glu Leu Leu Asn Pro Pro Gln Lys Ser Glu 100 105 110

Ser Gly Gly Pro Val Arg Tyr Gly Ala Arg Arg Gln Arg Leu Pro Glu 115 120 125

Met Leu Thr Arg Ile Val Gly Ser Val Thr Ala Ala Arg Ser Lys His 130 135 140

Leu Leu Gly Gly Val Leu Gln Pro Thr Gly Lys Trp Ala Phe Phe 145 \$150\$

Val Ser

<210> 137

<211> 46 <212> PRT

<212> PRT <213> Homo sapien

<400> 137

Met Arg Ile Cys Leu Cys Phe His Leu Leu Ser Leu Cys Leu Cys Leu 1 5 10 15

Ser Asn Leu Pro Pro Gln Glu His Gln Gly Asn Gln Cys Cys Arg Pro 20 25 30

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Arg Leu Asn Leu Arg Phe Leu Asn Gly Arg Asn Glu Ser Thr
                         40
<210> 138
<211> 47
<212> PRT
<213> Homo sapien
<400> 138
Met Leu Phe Gln Gln Leu Leu Lys Leu Phe Cys Gln Met Leu Phe Leu
1 5
Gln Cys Glu Gln Asn Ile Asn Arg Ile Lys Asn Ile Asn Arg Thr Glu
          20
                             25
Asn Lvs Asn His Val Ile Ile Ser Ile Leu Ile Gly Leu His Met
                         40
<210> 139
<211> 68
<212> PRT
<213> Homo sapien
<400> 139
Met Leu Ala Gly Ser Leu Leu Leu Ala Phe Glu Arq Gly Ala Ala Val
Ser Ser Val Cys Arg Val Lys Lys Arg Gly Gln Ser Ser Glu Glu Met
               25
Asn Pro Ser Arg Lys Gly Cys Leu Thr Ser Arg Arg Glu Asp Asn Gln
                         40
Glu Gly Leu Trp Val Ser Leu Leu Ser Tyr Thr Ser Phe Gln Val Thr
    50 55
Trp Asp His Gly
65
<210> 140
<211> 13
<212> PRT
<213> Homo sapien
<400> 140
Met Trp Gln Val Glu Asp Gly Phe Leu Ser Ser Leu Thr
             5
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101

<210> 141 <211> 108 <212> PRT <213> Homo sapien

<400> 141

Met Gln Pro Arg Leu Phe Ser Cys Ala Tyr Val Ser Arg Met Pro Thr

Phe Leu Phe Asn Ala Met Gln Ile Ser Lys Lys Ser Leu Thr Met Lys

Ala Ala Cys Arg Gln Lys Ser Phe Gln Cys Ile His Val His Ile His 35 40

Thr Leu His Arg His Met Ser Ser Tyr Lys Glu Ile Thr Gln Asp Gln

Asn Phe Thr Tyr Cys Gln Phe Leu Lys Val Gln Pro His Phe Gln Lys 70

Pro Gln Thr Ser Ile Arg Ser Asn Glu Ile Tyr Ile Phe Leu Ile Leu

Gly Lys Cys Asn Ile Pro Val Leu Glu Gln Gly Glu

<210> 142 <211> 38 <212> PRT <213> Homo sapien

<400> 142

Met Ser Leu Leu Ile Leu Thr Met Lys Leu Lys Lys Lys His Leu 5

Lys Ser Thr Glu Ile Arg Val Lys Val Thr Val Ile Thr Tyr Leu Tyr 20 25

His Asn Ile Leu Val Asp 35

<210> 143

<211> 74 <212> PRT <213> Homo sapien

<400> 143

102

Met His Thr His His Tyr Tyr Gly Val Ser Tyr Pro Gln Leu Ser Pro

Asp Gln Ala Leu Lys Ala Gly Arg Ala Arg Ser Gly Ile Pro Gly Lys $20 \\ 25 \\ 30$

Gly Trp Glu Gly Leu Ala Leu Arg Lys Gly Cys Glu Thr Gly Met Arg 35 40 45

Trp Gly Glu Cys Ser Glu Gly Gly Gly Lys Gly Val Pro Ala Gly Gly 50 55 60

Val Cys Ser Ser Thr Ala Glu Ala Ala Glu 65 70

<210> 144 <211> 119 <212> PRT

<213> Homo sapien

<400> 144

Met Ile Ala Arg Gly Leu Ala Cys Cys Leu Leu Asp Ser Phe Leu Leu 1 5 10 15

Leu Phe Ser Leu Pro Val Gly Trp Thr Cys His Cys Cys Thr Cys Ala 20 25 30

Phe Ala Phe Ser Tyr Ser Phe Phe His Leu Leu Leu Ser Ile Cys Asp 35 40 45

Thr Ser Trp Cys Val Ser Tyr Arg Trp Pro Ser Ser Cys Cys Arg Ser 50 60

Leu Ala Leu Pro Gly Val Ser Ser Leu Ser Arg Val Pro Pro Leu Leu 65 707580

Pro Ser Cys Arg Leu Arg Phe Gly Gly Pro Ser Val Arg Val Arg Phe 85 90 95

Pro Ile Val Pro Gly Tyr Pro Met Trp Ala Pro Leu Ala Arg Ser Pro $100 \hspace{1cm} 105 \hspace{1cm} 110 \hspace{1cm}$

Pro Phe Gly Asn Arg Phe Arg 115

<210> 145

103

<211> 23

<212> PRT <213> Homo sapien

<400> 145

Met Ile Lys Gly Ile Gly Lys Ser Thr Lys Thr Lys Ser Ser Asp Glu

Thr Gln Glu Ser Gly Arg Arg 20

<210> 146 <211> 33 <212> PRT <213> Homo sapien

<400> 146

Met Leu Asn Leu Leu Ile Ile Ser Pro Leu Asp Cvs Lvs Ile Tvr Val 5

Gly Arg Asp Lys Ile Val Ser Val Leu Ile Val Ser Pro Lys Pro Leu

Glu

<210> 147

<211> 130 <212> PRT

<213> Homo sapien

<400> 147

Met Gln Leu Gln Ala Pro Trp Pro Gln Cys Ser Ser Lys His Gln Val

Cys Thr Cys Leu Gly Gln Ser Val Leu Gly Ile Pro Ser Ala Leu Asn

Val Val Leu Pro Glu Lys Cys Ile Val His Ser Tyr Ile Leu Lys Val

Ser Leu His Cys Tyr Leu Ser Arg Ser Ser Leu Asn Ile Tyr Ile Ser

Ile Pro Phe Pro Pro Cys Phe Met Tyr Val His Ser Thr His His Asp 70 75

Leu Thr Leu Leu Ser Ile Tyr Ser Phe Thr Val Cys Phe Pro Ile Pro

> 104 90

85

95

Lys Tyr Lys Leu Ser Lys Asp Arg Asn Phe Cys Ala Leu Leu Leu Asn 100

Leu Gln Phe Val Glu Lys Cys Leu Pro Tyr Ile Lys His Ser Val Asn 120

Val Tyr 130

<210> 148 <211> 51

<212> PRT <213> Homo sapien

<400> 148

Met Ser Ser Lys Glu Val Ser Leu Asp Ser Leu Leu Leu Gly Arg Leu

Met Arg Gly Tyr Gln Arg Thr Glu Phe Asn Cys Leu Trp Ile Leu Leu 20

Lys Thr Leu Arg Ala Ser Gln Gly Ala Tyr Met Pro Arg Leu Ser Leu

Gly Ile Gly

<210> 149

<211> 42 <212> PRT

<213> Homo sapien

<400> 149

Met Ala Ile Ser Leu Ile Val Cys Lys Ile Tyr Lys Ala Tyr Leu Asn

Lys Phe Phe Leu Phe Ile Val Met Lys Thr Arq Ser Ser Leu Ser Gln

His Thr Ser Ser Asn Tyr Leu Gln Pro Leu

<210> 150 <211> 44

<212> PRT <213> Homo sapien

105

<400> 150

Met Ser Glu Arg Ala Pro Lys Thr Phe Ser Cys Phe Phe Val Phe Phe

Tyr Pro Thr Trp His Thr Trp His Gln Pro Asn Cys Phe Ile Thr Glu

Asn Lys Tyr Phe Leu Pro Lys Tyr Leu Phe His Arg 4.0

<210> 151

<211> 68 <212> PRT

<213> Homo sapien

· <400> 151

Met Tyr Ile Phe Phe Gly Ser Ala Val Leu Leu Arg Ile Phe His

Pro Thr Gly Tyr Lys Lys Met Phe Thr Ile Val Leu Phe Leu Leu Leu

Glu Ser Gly Thr Gln Tyr Lys Gln Gln Ser Leu Arg Asp Trp Ser His

Val Ser Thr Gln Glu His Arg Asp Gly Ile Gln Arg Leu Ile Ile Met

Trp Leu Ser Leu 65

<210> 152 <211> 91 <212> PRT <213> Homo sapien

<400> 152

Met Trp Ser Ile His Ser Leu Lys Leu Thr Leu Ser Glu Thr Thr Asp 5

Ser Arg Ser Ser Asn Asn Ala Thr Ala Phe Asn Met Val Ser Leu Ile 25

Thr Leu His Asp His Lys Glu Ala Ala Ile Tyr Ser Phe Cys Leu Arg 35 40

106

Leu Tyr Leu Gln Glu Pro Ile Asp Lys Val Asn Cys Phe Tyr Phe Lys

Leu Gln Ser Val Glu Tyr His His Ser Gln Ser Lys Leu His Glu Asn

Arg Ala Tyr Ile Phe Val Thr His Glu Gly Gly 85

<210> 153 <211> 156 <212> PRT <213> Homo sapien

<400> 153

Met His Arg Ser Gly His Leu Ser Trp Val Thr Ser Leu Val Ala Ile 10

Gly Leu Gly Ser Gly Ala Ala Cys Thr Glu Arg Arg Pro Trp Ala Gln

Arg Ala Ala Gly Ala Pro Gly Gly Ser Phe Thr Glu Ala Gly Leu Gly

Leu Ala Pro Glu Pro Val Arg Ser Gly Val Gly Ser Gly Ala Gly Asp 55

Ser Ala Met Ala Thr Ala Gly Ala Ala Cys Val Ser Ala Val Ala Pro 70 65 75

Ser Asp Pro Ile Gly Ala Arg Asp Leu Ser Gly Ala Ala Ala Ala Gly 85

Pro Gly Arg Thr Gly Glu Gly Asp Ala Gly Arg His Gly Asp Leu Gly 100

Arg Arg Ala Gly Gly Gly His Ala Gly Gly Ser Ala Gly Arg Gly 115 120

Arg Gly Gly His Arg Ala Ala Glu Lys Val Pro Met Ala Trp Gly Ser

Gly Ala Arg Val Gly Lys Thr Gln Asp Ile Ala Ser 145 150

<210> 154 <211> 22

107

<212> PRT <213> Homo sapien

<400> 154

Met Phe Arg Asp Leu Gly Tyr Phe Lys Glu Val Ser Ala Ala Val Val

Thr Asp Val Glu Leu Leu

<210> 155

<211> 46 <212> PRT <213> Homo sapien

<400> 155

Met Gly Glu Glu Leu Gly Arg Gln Glu Thr Ala Tyr Asn Leu Thr Asp

Gln Glu Lys Gly Ser Gly Phe Tyr Ser Lys Cys Lys Gly Lys Leu Trp 25

Glu Arg Phe Lys Gln Ile Ser Lys Thr Ile Phe Tyr Leu Gln 40

<210> 156

<211> 51 <212> PRT

<213> Homo sapien

<400> 156

Met Tyr Thr Ser Asn Gln Tyr Ile Tyr Leu Lys Leu Trp Leu Glu Leu 10 15

Thr Ala Leu Met Asn Thr Met Cys Pro Gln Lys Leu Thr Ile Glu Gly 20

Phe Lys Thr Lys Lys Leu His Thr Thr Thr Phe Leu His Thr Leu Asn

Lys Lys Ile 50

<210> 157

<211> 72 <212> PRT <213> Homo sapien

<400> 157

108

Met Ala Ser Leu Asn Ser Lys Ile Leu Pro Ser Ser Lys Leu Leu Leu Gly Leu Met Val Lys Ile Leu Thr Asn Gly Ser Lys Ile Pro Tyr Leu 20 25 Pro His Pro Pro Ile Pro Thr Val Leu Arg Pro Leu Ser Pro Arg Ser Trp Ile Leu Ser Ser Gln Val Val Ser Ala Cys His Gln Lys Glu Cys Asn Ser Val Leu Asp Leu His Thr 65 <210> 158 <211> 32 <212> PRT <213> Homo sapien <400> 158 Met Lys Cys Arg Gln Met Ala Arg Cys Cys Glu Val Lys Trp Leu Trp His Leu Val Ala Thr Glu Thr Thr Val Ile Lys Ile Met Ile Ile Thr <210> 159 <211> 19 <212> PRT <213> Homo sapien <400> 159 Met Ser Leu Val Ser Gly Glu His Lys Ile Ile Asp Ser Ala Glu Val 5 Trp Ser Lvs <210> 160 <211> 46 <212> PRT <213> Homo sapien <400> 160 Met Tyr Thr Glu Ile Ser Lys Val Leu Gln Lys Tyr Pro Lys Ser Arg 5 10

109

Val Ser Ile Ser His Phe Thr Ser Ser Ala Cys Phe Thr Pro Ile Leu 25

Asp Cys Phe Ile Ser Glu Leu Asp Val Ile Pro Arg Val Arg

<210> 161

<211> 47 <212> PRT

<213> Homo sapien

<400> 161

Met Pro Gln Trp His Gln Pro Ser His Leu Glu Lys Lys Ser Leu Ser

Ser Val Met Cys Ser Ala Arg Pro Pro Met Val Gln Thr Tyr Cys Thr 20

Cys Leu Val Ser Pro Ala Leu Ala His Leu Pro Leu Tyr Ser Leu 4.0

<210> 162

<211> 137 <212> PRT

<213> Homo sapien

<400> 162

Met Ser Gln Asp Thr Arg Ser Cys Pro Leu Phe Leu Ala Gln Leu Gly 5

Arg Arg Lys Gly Leu Gln Ala Arg Ala Ala Gly Gln Ala Gly Leu Pro

Leu Gly His Arg Thr Pro Leu Pro Pro Arg Pro Arg Leu His Ser His

His His Lys Ala Gln Val Pro Ser His Trp Leu Pro Lys Lys Ala Thr 50 55

Glu Arg Ile Phe Phe Leu Pro Leu Asn Val Ser Phe Pro Leu Gly Cys

Leu Ser Val Ala Leu Pro Ser Gln Val Phe Leu Gly Met Leu Arg Ala 85

Trp Arg Cys Thr Gly Gly Val Gln Trp His Leu Pro Pro Glu Leu Pro 100 105

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His Ser Leu Leu Arg Asn Leu Arg Gly Glu Gly Gly Ala Pro Gly Leu
Arg Glu Lys Glu Ser Val Leu Thr Phe
   130
                   135
<210> 163
<211> 68
<212> PRT
<213> Homo sapien
<400> 163
Met Pro Val Ile Pro Ala Leu Trp Glu Ala Glu Ala Gly Gly Ser Arg
Gly Gln Glu Ile Glu Thr Ile Leu Ala Asn Ala Val Lys Ser Arg Leu
Leu Lys Ile Gln Lys Ile Ser Gln Ala Trp Trp Arg Ala Pro Val Val
Pro Ala Thr Arg Glu Ala Glu Ala Gly Glu Trp Leu Glu Pro Gly Arg
Arg Ser Leu Gln
65
<210> 164
<210> 164
<211> 17
<212> PRT
<213> Homo sapien
<400> 164
Met Asn Ile Thr Ser Tyr Lys Ser Phe Ser Ile Tyr Ile Asn Ala Ile
                                      10
Asp
<210> 165
<211> 784
<212> PRT
<213> Homo sapien
<400> 165
Met Ala Ser Leu Asp Asp Pro Gly Glu Val Arg Glu Gly Phe Leu Cys
                                       10
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Pro Leu Cys Leu Lys Asp Leu Gln Ser Phe Tyr Gln Leu His Ser His 2.0 Tyr Glu Glu Glu His Ser Gly Glu Asp Arg Asp Val Lys Gly Gln Ile Lys Ser Leu Val Gln Lys Ala Lys Lys Ala Lys Asp Arg Leu Leu Lys 55 Arg Glu Gly Asp Asp Arg Ala Glu Ser Gly Thr Gln Gly Tyr Glu Ser 70 Phe Ser Tyr Gly Gly Val Asp Pro Tyr Met Trp Glu Pro Gln Glu Leu 90 85 Gly Ala Val Arg Ser His Leu Ser Asp Phe Lys Lys His Arg Ala Ala Arg Ile Asp His Tyr Val Val Glu Val Asn Lys Leu Ile Ile Arg Leu Glu Lys Leu Thr Ala Phe Asp Arg Thr Asn Thr Glu Ser Ala Lys Ile Arg Ala Ile Glu Lys Ser Val Val Pro Trp Val Asn Asp Gln Asp Val 145 150 155 Pro Phe Cys Pro Asp Cys Gly Asn Lys Phe Ser Ile Arg Asn Arg Arg 165 His His Cys Arg Leu Cys Gly Ser Ile Met Cys Lys Lys Cys Met Glu 185 Leu Ile Ser Leu Pro Leu Ala Asn Lys Leu Thr Ser Ala Ser Lys Glu 200 Ser Leu Ser Thr His Thr Ser Pro Ser Gln Ser Pro Asn Ser Val His Gly Ser Arg Arg Gly Ser Ile Ser Ser Met Ser Ser Val Ser Ser Val 225 230 235 Leu Asp Glu Lys Asp Asp Asp Arg Ile Arg Cys Cys Thr His Cys Lys 245 250 255

112

Asp Thr Leu Leu Lys Arg Glu Gln Gln Ile Asp Glu Lys Glu His Thr 260 265 Pro Asp Ile Val Lys Leu Tyr Glu Lys Leu Arg Leu Cys Met Glu Lys 275 280 Val Asp Gln Lys Ala Pro Glu Tyr Ile Arq Met Ala Ala Ser Leu Asn Ala Gly Glu Thr Thr Tyr Ser Leu Glu His Ala Ser Asp Leu Arg Val Glu Val Gln Lys Val Tyr Glu Leu Ile Asp Ala Leu Ser Lys Lys Ile 325 330 Leu Thr Leu Glv Leu Asn Gln Asp Pro Pro Pro His Pro Ser Asn Leu 345 Arg Leu Gln Arg Met Ile Arg Tyr Ser Ala Thr Leu Phe Val Gln Glu Lys Leu Leu Gly Leu Met Ser Leu Pro Thr Lys Glu Gln Phe Glu Glu Leu Lys Lys Lys Arg Lys Glu Glu Met Glu Arg Lys Arg Ala Val Glu Arg Gln Ala Ala Leu Glu Ser Gln Arg Arg Leu Glu Glu Arg Gln Ser 405 410 415 Gly Leu Ala Ser Arg Ala Ala Asn Gly Glu Val Ala Ser Leu Arg Arg 420 425 Gly Pro Ala Pro Leu Arg Lys Ala Glu Gly Trp Leu Pro Leu Ser Gly 435 440 Glv Gln Glv Gln Ser Glu Asp Ser Asp Pro Leu Leu Gln Gln Ile His 450 455 460 Asn Ile Thr Ser Phe Ile Arg Gln Ala Lys Ala Ala Gly Arg Met Asp 465 470 475 Glu Val Arg Thr Leu Gln Glu Asn Leu Arg Gln Leu Gln Asp Glu Tyr 485 490

113	
Asp Gln Gln Gln Thr Glu Lys Ala Ile Glu	Leu Ser Arg Arg Gln Ala
500 505	510
Glu Glu Asp Leu Gln Arg Glu Gln Leu	Gln Met Leu Arg Glu Arg
515 520	525
Glu Leu Glu Arg Glu Arg Glu Gln Phe Arg	Val Ala Ser Leu His Thr
530 535	540
Arg Thr Arg Ser Leu Asp Phe Arg Glu Ile	Gly Pro Phe Gln Leu Glu
545 550	555 560
Pro Ser Arg Glu Pro Arg Thr His Leu Ala 565 570	
Ser Ser Pro Val Pro Ser Ser Thr Ala Pro	Lys Thr Pro Ser Leu Ser
580 585	590
Ser Thr Gln Pro Thr Arg Val Trp Ser Gly 595 600	Pro Pro Ala Val Gly Gln 605
Glu Arg Leu Pro Gln Ser Ser Met Pro Gln	Gln His Glu Gly Pro Ser
610 615	620
Leu Asn Pro Phe Asp Glu Glu Asp Leu Ser	Ser Pro Met Glu Glu Ala
625 630	635 640
Thr Thr Gly Pro Pro Ala Ala Gly Val Ser	Leu Asp Pro Ser Ala Arg
645 650	655
Ile Leu Lys Glu Tyr Asn Pro Phe Glu Glu	Glu Asp Glu Glu Glu Glu
660 665	670
Ala Val Ala Gly Asn Pro Phe Ile Gln Pro	Asp Ser Pro Ala Pro Asn
675 680	685
Pro Phe Ser Glu Glu Asp Glu His Pro Gln 690 695	Gln Arg Leu Ser Ser Pro
Leu Val Pro Gly Asn Pro Phe Glu Glu Pro	Thr Cys Ile Asn Pro Phe
705 710	715 720
Glu Met Asp Ser Asp Ser Gly Pro Glu Ala	Glu Glu Pro Ile Glu Glu
725 730	735
Glu Leu Leu Gln Gln Ile Asp Asn Ile	Lys Ala Tyr Ile Phe Asp

114

740 745 750

Ala Lys Gln Cys Gly Arg Leu Asp Glu Val Glu Val Leu Thr Glu Asn 755 760 765

Leu Arg Glu Leu Lys His Thr Leu Ala Lys Gln Lys Gly Gly Thr Asp

<210> 166

<211> 96 <212> PRT

<213> Homo sapien

<400> 166

Met Ser Gly Thr Ser Thr Pro Ala Met Gly Val Cys Ala Glu Pro Leu 1 5 10 15

Lys Val Asp Leu Ser Phe Gly Glu Pro Ser Glu Arg His Ser Trp Tyr 20 25 30

Leu Trp Glu Trp Leu Trp Gly Val Leu Trp His His Thr Asp Asn Phe 35 40 45

Val Phe Leu Ile Gly Ile His Gly Leu Gly Ser Trp Asp Gly Gly Arg $50 \hspace{1.5cm} 60$

Gly Lys Pro Gln Ser Pro Trp Lys Cys Met Gln Asn Ile Trp Val Cys 65 70 75 80

Arg Cys Ile Ile Leu Arg Leu Trp Pro Leu Gln Phe Phe Gln Arg Val

<210> 167

<211> 215

<212> PRT

<213> Homo sapien

<400> 167

Met Ser Gly Glu Lys Arg Ser Pro Ser Val Asn Thr Gln Ala Pro Gly
1 5 10 15

Thr Thr Arg Gly Ala Cys Lys Gly Thr Thr Ser Trp Glu Pro Ser Trp
20 25 30

Pro His Cys Glu Glu Val Pro Gln Glu Glu Lys Phe Leu Arg Gly Thr 35 40 45

115

Phe Phe Phe Gln Phe Leu Ala Ala Gly Gly Trp Gly Ser Ser Ser Val

Val Ser Arg Arg Val Arg Pro Arg Gln Arg Pro Thr Ser Arg Arg Arg 115 120 125

Pro Gly Val Glu Ser Ala Leu Asn Lys Ile Leu Ser Gln Leu Cys Val 130 135 140

Pro Arg Arg Val Lys Gly Pro Val Pro Ala Gly Ala Arg Gly Ser Ala 145 155 160

Trp Ser Trp Arg Pro Ala Arg Arg Leu Trp Arg Pro Leu Ser Glu Pro

Arg Arg Pro Pro Arg Val Phe Val Asp Cys Arg Ser Pro Gly Arg Leu 180 $$185\mathcharpoons$

Gly Thr Val Thr Ala Pro Lys Ala Gly Asp Val Ala Ala Leu Gly Arg 195 200 205

Arg Leu Leu Pro Pro Pro Leu 210 215

<210> 168 <211> 67 <212> PRT <213> Homo sapien <400> 168

Met Ser Leu arg Pro Leu Met Gly Leu asp Pro Ala Ser Gly Lys Leu 1 5 5 10 15

Leu Ser His Thr Leu Pro His Leu Thr Ile His Val Ser Phe Leu Pro

Leu Ser Leu His Ala Cys His Ser His Ser Leu Cys Met Pro Ala Phe 35 40 45

116

Pro Ala Ala Pro Cys Thr Leu Arg Leu Trp Pro Phe Pro Ala Phe Cys 50 55 Leu Pro Gly <210> 169 <211> 102 <212> PRT <213> Homo sapien <400> 169 Met Pro Thr Pro Ser Leu Ser Tyr Thr Gly Glu Gly Gln Lys Gln Asp Val Glu Phe Ala Trp Pro Trp Pro Thr Gln Thr Thr Ser Pro Ser Ser Gly Ser Gln Ala Arg Glu Lys Gly Ile Ser Arg Thr Thr Val Pro Gly 40 Ile Lys His Lys Leu Lys Leu Ala Gly Ala Gly Met Val Ser Gln Thr Ser Glu Leu Pro Pro His Leu His Leu Phe Gly Leu Thr Gly Asn Met 75 Ala Ser Leu Gly Pro Ala Gly Arq Ala His Gly Arq Arq Leu Leu Ser 85 90 His Gln Ala Thr Glu Asp 100 <210> 170 <211> 38 <212> PRT <213> Homo sapien <400> 170

Met Arg Trp Val Glu Ser Thr Leu Glu Asn Ile Ile His Lys Ala Asn

Asn Gln Thr Gly Arg Lys Asp Asn Thr Lys Ser Leu Thr Ser Thr Ile \$20\$ \$25\$ 30

His Gly Ser Leu Ala Leu

117

35 <210> 171 <211> 52 <212> PRT <213> Homo sapien <400> 171 Met Ser Gln Phe Leu Asp Leu Leu Thr Val Ile Asn Ser Gln Ala His Ser Leu Ser Arg Val Asn Asp Thr Thr Phe Asn Leu Glu Lys Ser Ser 20 25 30 Ile Thr Phe Tyr His Phe Gln Ile Ser Phe Leu Pro Ser Leu His Thr Lys Asp Ser Ser 50 <210> 172 <211> 45 <212> PRT <213> Homo sapien <400> 172 Met Lys Leu His Met Thr Arg Phe Ser Phe Lys Leu Ile Leu Lys Cys

Leu Leu Leu Gly Val Leu Asn Ser Phe Thr Val Leu Val Thr Thr 2.0 25

Arg Leu Ile Ser Ala Ile Leu Tyr Tyr Ala Ile Phe Ser 40

<210> 173 <211> 34 <212> PRT <213> Homo sapien

<400> 173

Met Leu Thr Cys Cys Leu Asn Thr Glu Asp Ala Ile Leu Phe Ser Arg

Ala Leu Phe Leu His Gly Lys Ile Val Asp Pro Thr Ile Lys Cys Gly

Tyr Ser

118

<210> 174 <211> 29 <212> PRT <213> Homo sapien <400> 174 Met Phe Leu Arg Val Ala Gly Asn Ile Leu Leu Glu Asn Gln Met Asn Thr Gly Arg Tyr Gly Tyr Gln Glu Leu Arg Leu Ile Leu 20 <210> 175 <211> 21 <212> PRT <213> Homo sapien <400> 175 Met Asn Ile His Asn Pro Ser Ser Phe Leu Val Ser Leu Cys Asn Leu 10 Ala Ser Gln Asn Tvr 20 <210> 176 <211> 78 <212> PRT <213> Homo sapien <400> 176 Met Lys Ser Lys Ser Ser Leu Pro Ser Leu Gly Val Lys Leu His Asn Leu Ile Ser Ala Ser Ser Cys Cys Leu Tyr Phe Arg Lys Lys Gln 25 Thr Val Gly Phe Pro Cys Glu Ser Asp Phe Ser Leu Ser Ala Tyr Cys 40 Ala Ser Ala Phe Pro Cys Val Ser His Asp Leu Met Ala Ser Leu Thr

Pro Asn Cys Met Tyr Pro Ala Gln Ile Ser Ile Leu Pro Gln

70

<210> 177

119

<211> 37 <212> PRT <213> Homo sapien

<400> 177

Met Ser Leu Lys Arg Tyr Cys Leu Ala Leu Lys Ser Ala Ser Val Val 10

Lys Val Leu Gln Tyr Ile His Thr Glu Glu Lys Cys Gly Ser Glu Ser

Pro Trp Ala Ala Phe 35

<210> 178

<211> 21 <212> PRT <213> Homo sapien

<400> 178

Met Thr Lys Pro Thr Arg Leu Leu Pro Lys Asn Leu Ile Tyr Cys Pro 10

Arg Ser Arg Leu Ser 20

<210> 179 <211> 57 <212> PRT <213> Homo sapien

<400> 179

Met Cys Cys Val Ser Gly Gly Trp Lys Lys Leu Ile Gln Leu Trp Pro

Ser Ser Ile Met Val Arg Tyr His Tyr His Leu Gln Ile Ser Ser Lys

Ile Gly Gly Leu Pro Ser Trp Ala Leu Arg Ala Asn Ser Leu Gly Gln 40

Phe Phe Leu Leu His Ser Thr Ile Glu 50

<210> 180 <211> 23 <212> PRT <213> Homo sapien

120

<400> 180

Met Asn Phe Ile Leu Cys Pro Cys Phe Asp Ala Ser Thr Gln Phe Leu

Val Ile Ser Val Lys Tyr Asn 20

<210> 181 <211> 92 <212> PRT <213> Homo sapien

<400> 181

Met Arg Lys Lys Cys Arg Leu Thr Ser Pro Cys Trp Trp Arg Trp Arg

Arg Gly Gly Glu Gln Leu Leu Arg Ile Phe Val Leu Ile Phe Ser Cys 25

Ser Pro Pro Asp Asn Thr Val Gly Thr Cys Ile Leu Trp Gly Ile Phe 40

Ser Gly Lys Ser Lys Asp Cys Glu Trp Leu His Leu Ile Ser Thr Leu

Arg Asn Ala Glu Ala Cys Ser Ser Gln Val Leu Lys Asn Tyr Leu Glu

Lys Lys Asn Arg Pro Ile Lys Ser Thr Val Lys Arg 85 90

<210> 182 <211> 27 <212> PRT

<213> Homo sapien

<400> 182

Met Leu His Pro Thr Glu Met Gly Pro Gln Val Pro Asp Leu Pro Phe

Arg Arg Gly Arg Gln Val Gly Glu Gln Gln Met 20

<210> 183 <211> 166 <212> PRT <213> Homo sapien

121

<400> 183

Met Pro Leu Thr Pro Glu Pro Pro Ser Gly Arg Val Glu Gly Pro Pro 1 5 10 15

Ala Trp Glu Ala Ala Pro Trp Pro Ser Leu Pro Cys Gly Pro Cys Ile \$20\$

Pro Ile Met Leu Val Leu Ala Thr Leu Ala Ala Leu Phe Ile Leu Thr 35 40

Thr Ala Val Leu Ala Glu Arg Leu Phe Arg Arg Ala Leu Arg Pro Asp 50 55 60

Pro Ser His Arg Ala Pro Thr Leu Val Trp Arg Pro Gly Gly Glu Leu 65 70 75 80

Trp Ile Glu Pro Met Gly Thr Ala Arg Glu Arg Ser Glu Asp Trp Tyr 85 90 90 95

Gln Val Gly Thr Leu Glu Ala Arg Ala Thr Ala Pro Pro Ala Pro Ser 115 120 125

Ala Pro Asn Ser Ala Pro Ser Asn Leu Gly Pro Gln Thr Val Leu Glu 130 135 140

Val Pro Ala Arg Ser Thr Phe Trp Gly Pro Gln Pro Trp Glu Gly Arg 145 150155155

Pro Pro Pro Gln Ala Trp 165

<210> 184

<211> 80

<212> PRT

<213> Homo sapien

<400> 184

Met Leu Thr Ser His Phe Ile Leu Ile Pro Val Ile Phe Ser Leu Ghn 1 10 15

Tyr Gln Cys Leu Gly Ala Arg Lys Leu Cys Gln Cys Gln Trp Leu Trp 20 25 30

122

Arg Trp Gln Lys Lys Gly Gly Gln Pro Pro Gly Thr Ala Glu Ser Lys 40

Pro Asp Ser Gln Pro Gln Lys Val Gly Gln Asp Ala Ala Asn Ser Ser 55

Asn Pro Lys Lys Ala Ala Glu Ile Thr Val Ile Gln Gln Thr Tyr Phe 70

<210> 185 <211> 159 <212> PRT <213> Homo sapien

<400> 185

Met Asp Thr Ile Leu Val Phe Ser Leu Ile Ile Ala Ser Tyr Asp Ala

Asn Lys Lys Asp Leu Arg Asp Ser Ser Cys Arg Leu Glu Gln Leu Pro 25

Gly Ile Phe Pro Lys Asp Val Arg Ser Ile Arg Glu Leu Gln Met Gln 40

Glu Thr His Thr Glu Thr Lys Arg Thr Thr Phe Ile Gln Asn Arg Thr

Ile Ala Thr Leu Gln Cys Leu Gly Ser Asp Ser Lys Val Lys Val Asn 70 75

Leu Val Tyr Leu Glu Arg Arg Pro Lys Val Lys His Ile Leu Lys Asn

Leu Arg Ile Ile Ala Ala Pro Arg Arg Asn Ser Ser Ala Ser Ser Ser 100 105

Cys His Leu Ile Pro Thr Ser Lys Phe Gln Thr Gly Ser Leu Leu Lys 115 120

Gly Lys Val Ser Met Pro Arg Ser Gln Glu Ala Val Pro Met Pro Val 130 135

Val Val Glu Met Ala Lys Glu Gly Arg Pro Ala Thr Trp Asp Ser 150

<210> 186 <211> 928

123

<212> PRT <213> Homo sapien

<400> 186

Met Ala Glu Gly Lys Glu Lys Gln Val Thr Ser Tyr Met Asp Gly Ser

Arg Pro Tyr Asp Val Ser Met Thr Tyr Ile His Lys Ala Gly Gly Pro 20 25 30

Asp Gln Gln Glu Leu Val Met Leu Thr Cys Thr Val Pro Leu Asp Ser 35 40 40 45

Cys Cys His Leu Pro Gln Ala Arg Thr Asn Tyr Arg Lys Tyr Phe Arg 50 55 60

Ser Glu Ala Ala Phe Thr Leu Ala Asp Phe Ile Tyr Lys Ser Met Ile 65 70 75 80

Arg Val Asn Ser Ser Arg Leu Val Arg Val Thr Gln Val Glu Asn Glu 85 90 95

Glu Lys Leu Lys Glu Leu Glu Glu Phe Ser Ile Trp Asn Phe Phe Ser 100 $$105\$

Ser Phe Leu Lys Glu Lys Leu Asn Asp Thr Tyr Val Asn Val Gly Leu 115 $$\rm 120$$

Tyr Ser Thr Lys Thr Cys Leu Lys Val Glu Ile Ile Glu Lys Asp Thr 130 135 140

Lys Tyr Ser Val Ile Val Ile Arg Arg Ser Trp Asp Val Ile Arg Val 145 150 155 160

Leu Lys Glu Leu Glu Gln Phe Ser Ile Trp Asn Phe Phe Ser Ser Phe 180 $$185\$

Leu Lys Glu Lys Leu Asn Asp Thr Tyr Val Asn Val Gly Leu Tyr Ser 195 200 205

Thr Lys Thr Cys Leu Lys Val Glu Ile Ile Glu Lys Asp Thr Lys Tyr 210 215 220

Ser Va 225	al Ile	val	Ile	Arg 230	Arg	Ser	Trp	Asp	Val 235	Ile	Arg	Val	Asn	Ser 240
Ser A	rg Let	Val	Arg 245	Val	Thr	Gln	Val	Glu 250	Asn	Glu	Glu	Lys	Leu 255	Lys
Glu Le	eu Gli	Gln 260	Phe	Ser	Ile	Trp	Asn 265	Phe	Phe	Ser	Ser	Phe 270	Leu	Lys
Glu Ly	ys Let 275		Asp	Thr	Tyr	Val 280	Asn	Gly	Ile	Pro	Trp 285	Thr	Lys	Val
Asp Ty	yr Phe	asp	Asn	Gly	Ile 295	Ile	СЛв	Lys	Leu	Ile 300	Glu	His	Asn	Gln
Arg Gl 305	ly Ile	Leu	Ala	Met 310	Leu	Авр	Glu	Glu	Сув 315	Leu	Arg	Pro	Gly	Val 320
Val Se	er Ası	Ser	Thr 325	Phe	Leu	Ala	Lys	Leu 330	Asn	Gln	Leu	Phe	Ser 335	Lys
His G	ly His	340	Glu	Ser	Lys	Val	Thr 345	Gln	Asn	Ala	Gln	Arg 350	Gln	Tyr
Asp Hi	is Thi		Gly	Leu	Ser	Cys 360	Phe	Arg	Ile	Cys	His 365	Tyr	Ala	Gly
Lys Va	al Thi	Tyr	Asn	Val	Thr 375	Ser	Phe	Ile	Asp	380	Asn	Asn	Asp	Leu
Leu Pl 385	ne Arg	Asp	Leu	Leu 390	Gln	Ala	Met	Trp	Lys 395	Ala	Gln	His	Pro	Leu 400
Leu Ar	rg Sei	Leu	Phe 405	Pro	Glu	Gly	Asn	Pro 410	Lys	Gln	Ala	Ser	Leu 415	Lys
Arg Pi	ro Pro	420	Ala	Gly	Ala	Gln	Phe 425	Lys	Ser	Ser	Val	Ala 430	Ile	Leu
Met Ly	ys Ası 435		Tyr	Ser	Lys	Ser 440	Pro	Asn	Tyr	Ile	Arg 445	Сув	Ile	Гуз
Pro As		His	Gln	Gln	Arg 455	Gly	Gln	Phe	Ser	Ser 460	Asp	Leu	Val	Ala
Thr G	ln Ala	Arg	Tyr	Leu	Gly	Leu	Leu	Glu	Asn	Val	Arg	Val	Arg	Arg

								1	25						
465					470					475					480
Ala	Gly	Tyr	Ala	His 485	Arg	Gln	Gly	Tyr	Gly 490	Pro	Phe	Leu	Glu	Arg 495	Tyr
Arg	Leu	Leu	Ser 500	Arg	Ser	Thr	Trp	Pro 505	His	Trp	Asn	Gly	Gly 510	Asp	Arg
Glu	Gly	Val 515	Glu	Lys	Val	Leu	Gly 520	Glu	Leu	Ser	Met	Ser 525	Ser	Gly	Glu
Leu	Ala 530	Phe	Gly	Lys	Thr	Lys 535	Ile	Phe	Ile	Arg	Ser 540	Pro	Lys	Thr	Leu
Phe 545	туг	Leu	Glu	Glu	Gln 550	Arg	Arg	Leu	Arg	Leu 555	Gln	Gln	Leu	Ala	Thr 560
Leu	Ile	Gln	Lys	Ile 565	Tyr	Arg	Gly	Trp	Arg 570	Сув	Arg	Thr	His	Tyr 575	Gln
Leu	Met	Arg	Lys 580	Ser	Gln	Ile	Leu	Ile 585	Ser	Ser	Trp	Phe	Arg 590	Gly	Asn
Met	Ala	Arg 595	Lys	Asn	Tyr	Arg	Lys 600	Tyr	Phe	Arg	Ser	Glu 605	Ala	Ala	Leu
Thr	Leu 610	Ala	Asp	Phe	Ile	Tyr 615	Lys	Ser	Met	Val	Gln 620	Lys	Phe	Leu	Leu
Gly 625	Leu	Lys	Asn	Asn	Leu 630	Pro	Ser	Thr	Asn	Val 635	Leu	Asp	Lys	Thr	Trp 640
Pro	Ala	Ala	Pro	Tyr 645	Lys	Сув	Leu	Ser	Thr 650	Ala	Asn	Gln	Glu	Leu 655	Gln
Gln	Leu	Phe	Tyr 660	Gln	Trp	Lys	Ala	Thr 665	Pro	Val	Pro	Pro	Ser 670	Ser	Gln
Сув	Lys	Arg 675	Phe	Arg	Asp	Gln	Leu 680	Ser	Pro	Lys	Gln	Val 685	Glu	Ile	Leu
Arg	G1u 690	Lys	Leu	Cys	Ala	Ser 695	Glu	Leu	Phe	Lys	Gly 700	Lys	Lys	Ala	Ser
Tyr 705	Pro	Gln	Ser	Val	Pro 710	Ile	Pro	Phe	Сув	Gly 715	Авр	Tyr	Ile	Gly	Leu 720

126

Gln Gly Asn Pro Lys Leu Gln Lys Leu Lys Gly Gly Glu Glu Gly Pro Val Leu Met Ala Glu Ala Val Lys Lys Val Asn Arg Gly Asn Gly Lys 740 Thr Ser Ser Arg Ile Leu Leu Thr Lys Gly His Val Ile Leu Thr Asp Thr Lys Lys Ser Gln Ala Lys Ile Val Ile Gly Leu Asp Asn Val 775 770 780 Ala Gly Val Ser Val Thr Ser Leu Lys Asp Gly Leu Phe Ser Leu His 785 Leu Ser Glu Met Ser Ser Val Gly Ser Lys Gly Asp Phe Leu Leu Val 805 810 Ser Glu His Val Ile Glu Leu Leu Thr Lvs Met Tvr Arg Ala Val Leu 820 825 Asp Ala Thr Gln Arg Gln Leu Thr Val Thr Val Thr Glu Lys Phe Ser Val Arg Phe Lys Glu Asn Ser Val Ala Val Lys Val Val Gln Gly Pro Ala Gly Gly Asp Asn Ser Lys Leu Arg Tyr Lys Lys Lys Gly Ser His 865 870 975 Cys Leu Glu Val Thr Val Gln Gln Leu Thr Ala Gly Tyr His Ala Gly 885 890 Gln Gly Glu Leu Ile Asn Phe Ser Ser Cys Leu Gln Ile Asn Leu Leu 905 Ser Glu His Lys Pro Arg Ala Ser Gly Thr Pro Cys Phe Glu Leu Arg <210> 187 <211> 96 <212> PRT <213> Homo sapien

<400> 187

127 Met Arg Arg Cys Tyr Ser Ile Pro Val Cys Lys Cys Ala Gly Met Pro Ala Leu Ser Asp Gly Gly His Asp Asn Met Ala His Ala Phe Lys Leu Thr Ser Asn Cvs Phe Trp Thr Thr Phe Asn Arg Gly Ser His Tyr His Gly Phe Lys Glu Pro Cys Gln Pro Arg Lys His Leu Thr Ala Gly Thr 55 Ala Gly Trp Ser Cys Cys Trp Leu Glu Val Tyr Ala Arg Ile Ala Lys 65 70 75 80 Asp Ser Trp Arg Met Gly Ser Pro Tyr Leu Cys Arg Leu Ala Ala Leu <210> 188 <211> 47 <212> PRT <213> Homo sapien <400> 188 Met Tyr Gln Lys Asp Leu Tyr Ile Ser Gln Arg Gly Thr Gln Ala Lys Leu Lys Ile Tyr Lys His Asn Gln Phe Thr Arg Glu Ile Ile Leu Thr 20 25 Val Phe Phe Leu Phe Phe Gln Thr Leu Leu Phe His Gly Lys Lys <210> 189 <211> 644 <212> PRT <213> Homo sapien <400> 189 Met Met Ile Ile Ala Leu Glu Phe Pro His Leu Val Val Asp Leu Ala Asp Asn Asn Trp Gln Cys Asp Asp Ser Val Ala Val Phe Gln Asn Phe

Ile Ser Glu Ser Trp Arg Lys Lys Trp Asn Val Ile Cys Asn Arg Ser

Ile	Gly 50	Ser	Glu	Glu	Ala	Asn 55	Gly	Gly	Thr	Pro	Gln 60	Ser	Arg	Ile	Ser
Arg 65	Glu	Thr	Arg	Leu	Pro 70	Pro	Ile	His	Leu	His 75	Arg	Met	Lys	Ser	Leu 80
Ile	Arg	Ser	Lys	Ala 85	Glu	Arg	Pro	Gln	Gly 90	Gly	Arg	His	Thr	Gly 95	Ile
Ser	Thr	Leu	Gly 100	Lys	Lys	Ala	Lys	Ala 105	Gly	Ser	Gly	Leu	Arg 110	Lys	Lys
Gln	Arg	Arg 115	Leu	Pro	Arg	Ser	Val 120	Arg	Ser	Thr	Arg	Asp 125	Val	Gln	Ala
Ala	Gly 130	Lys	Lys	Glu	Asp	Ala 135	Pro	Gln	Asp	Leu	Ala 140	Leu	Ala	Val	Cys
Leu 145	Ser	Val	Phe	Ile	Thr 150	Phe	Leu	Val	Ala	Phe 155	Ser	Leu	Gly	Ala	Phe 160
Thr	Arg	Pro	Tyr	Val 165	Asp	Arg	Leu	Trp	Gln 170	Lys	Lys	Cys	Gln	Ser 175	Lys
Ser	Pro	Gly	Leu 180	Asp	Asn	Ala	Tyr	Ser 185	Asn	Glu	Gly	Phe	Туг 190	Asp	Asp
Met	Glu	Ala 195	Ala	Gly	His	Thr	Pro 200	His	Pro	Glu	Thr	His 205	Leu	Arg	Gln
Val	Phe 210	Pro	His	Leu	Ser	Leu 215	Tyr	Glu	Asn	Gln	Thr 220	Pro	Phe	Trp	Val
Thr 225	Gln	Pro	His	Pro	His 230	Ala	Thr	Val	Ile	Pro 235	Asp	Arg	Thr	Leu	Gly 240
Arg	Ser	Arg	Lys	Asp 245	Pro	Gly	Ser	Ser	Gln 250	Ser	Pro	Gly	Gln	Сув 255	Gly
Asp	Asn	Thr	Gly 260	Ala	Gly	Ser	Gly	Asn 265	Asp	Gly	Ala	Val	Tyr 270	Ser	Ile
Leu	Gln	Arg 275	His	Pro	His	Ala	Gly 280	Asn	Arg	Glu	Leu	Met 285	Ser	Ala	Ala

Gln	Asp 290	His	Ile	His	Arg	Asn 295	Asp	Ile	Leu	Gly	Glu 300	Trp	Thr	Tyr	Glu
Thr 305	Val	Ala	Gln	Glu	Glu 310	Pro	Leu	Ser	Ala	His 315	Ser	Val	Gly	Val	Ser 320
Ser	Val	Ala	Gly	Thr 325	Ser	His	Ala	Val	Ser 330	Gly	Ser	Ser	Arg	Tyr 335	Asp
Ser	Asn	Glu	Leu 340	Asp	Leu	Pro	Leu	Ser 345	Gly	Glu	Ile	Thr	Ala 350	Ser	Leu
Cys	Lys	Met 355	Leu	Thr	His	Ala	Glu 360	Ala	Gln	Arg	Thr	Gly 365	Asp	Ser	Lys
Glu	Arg 370	Gly	Gly	Thr	Glu	Gln 375	Ser	Leu	Trp	Авр	Ser 380	Gln	Met	Glu	Phe
Ser 385	Lys	Glu	Arg	Gln	Val 390	Ser	Ser	Ser	Ile	Asp 395	Leu	Leu	Ser	Ile	Gln 400
Gln	Pro	Arg	Leu	Ser 405	Gly	Ala	Arg	Ala	Glu 410	Glu	Ala	Leu	Ser	Ala 415	His
Tyr	Ser	Glu	Val 420	Pro	Tyr	Gly	Asp	Pro 425	Arg	Asp	Thr	Gly	Pro 430	Ser	Val
Phe	Pro	Pro 435	Arg	Trp	Asp	Ser	Gly 440	Leu	Asp	Val	Thr	Pro 445	Ala	Asn	Lys
Glu	Pro 450	Val	Gln	Lys	Ser	Thr 455	Pro	Ser	Asp	Thr	Сув 460	Cys	Glu	Leu	Glu
Ser 465	Asp	Cys	Asp	Ser	Asp 470	Glu	Gly	Ser	Leu	Phe 475	Thr	Leu	Ser	ser	Ile 480
Ser	Ser	Glu	Ser	Ala 485	Arg	Ser	Lys	Thr	Glu 490	Glu	Ala	Val	Pro	Asp 495	Glu
Glu	Ser	Leu	Gln 500	Asp	Glu	Ser	Ser	Gly 505	Ala	Ser	Lys	Asp	Asn 510	Val	Thr
Ala	Val	Asp 515	Ser	Leu	Glu	Glu	Asn 520	Val	Thr	Phe	Gln	Thr 525	Ile	Pro	Gly
Lys	Cys	Lys	Asn	Gln	Glu	Asp	Pro	Phe	Glu	Lys	Pro	Leu	Ile	Ser	Ala

540

130 535

Pro Asp Ser Gly Met Tyr Lys Thr His Leu Glu Asn Ala Ser Asp Thr

545 550 555

Asp Arg Ser Glu Gly Leu Ser Pro Trp Pro Arg Ser Pro Gly Asn Ser 570

Pro Leu Gly Asp Glu Phe Pro Gly Met Phe Thr Tyr Asp Tyr Asp Thr 585

Ala Leu Gln Ser Lys Ala Ala Glu Trp His Cys Ser Leu Arg Asp Leu 600

Glu Phe Ser Asn Val Asp Val Leu Gln Gln Thr Pro Pro Cys Ser Ala 610 615 620

Glu Val Pro Ser Asp Pro Asp Lys Ala Ala Phe His Glu Arg Asp Ser 625 630 635

Asp Ile Leu Lys

530

<210> 190

<211> 48 <212> PRT

<213> Homo sapien

<400> 190

Met Trp Thr Phe Tyr Ser Lys His His His Val Leu Leu Lys Phe Pro

Gln Ile Leu Val Asp Val Leu Gln Gln Thr Pro Pro Cys Ser Ala Glu

Val Pro Ser Asp Pro Asp Lys Ala Ala Phe His Glu Arg Phe Leu Phe 40